# Smooth Emulator & Simplex Sampler User Manual

A BAND Collaboration Project

https://bandframework.github.io



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# 1 Overview

This manual describes how to install and run  $Smooth\ Emulator$  software. The software performs three basic functions. First, the  $Simplex\ Sampler$  chooses a set of points in model parameter space, at which full model runs will performed to then tune the emulator. The user must provide a description of the model parameters and the prior in a text files in a standard format. There are several options, the first of which is to choose the points that represent a simplex, e.g. an equilateral triangle in two dimensions or a tetrahedron in three dimensions. In a simplex, all points are equidistant from one another, and the number of training points is  $N_p + 1$ , where  $N_p$  is the number of parameters. In addition to the standard simplex, there are additional options which are motivated by the simplex form. For the standard form the  $N_p + 1$  training points in the simplex match the number of points needed to determine a linear fit. Another choice, which is based on the simplex chooses enough points to determine a quadratic fit,  $(N_p + 1)(N_p + 2)/2$ . The software will write the information about the training points in a standard format, which is described in the manual. If the user decides to use training points from a different procedure, the user can still record the information about the points in a same format, and the emulator tuning will still work, as the emulator itself is not predicated on a specific choice of training points.

The user is then responsible for running the full model at the training points and expressing observables, and the uncertainties, for each training point in a standard format. The manual describes the output format.

The second functionality of the software is to build and tune the emulator, referred here as *Smooth Emulator*. The emulator reads the information above, along with another user-provided parameter file to choose which observables are to be emulated, which parameters will be varied, and which emulator options will be applied. After being trained, the Taylor coefficients representing the emulator are written to a file. One can always add additional training points, and retrain the emulator.

The third functionality of the software is to perform a MCMC exploration of parameter space using the emulator. This user must express the experimental observables and their uncertainties in a standard format. The MCMC software will read the emulator coefficients from file and perform the MCMC exploration. This procedure is also guided by a simple text file of parameters. The MCMC software uses python and Matplotlib to generate plots that describe the posterior.

# 2 Installation and Getting Started

# 2.1 Prerequisites

Smooth Emulator software should run on UNIX, Mac OS or Linux, but is not supported for Windows OS. Smooth Emulator is largely written in C++. In addition to a C++ compiler, the user needs the following software installed.

- git
- CMake
- Eigen3 (Linear Algebra Package)
- Python/Matplotlib (only for generating plots in the MCMC procedure)

CMake is an open-source, cross-platform build system that helps automate the process of compiling and linking for software projects. Hopefully, CMake will perform the needed gymnastics to find the Eigen3 installation. To install CMake, either visit the CMake website (https://cmake.org/), or use the system's package manager for the specific system. For example, on Mac OS, if one uses homebrew as a package manager, the command is

#### % brew install cmake

Eigen is a C++ template library for vector and matrix math, i.e. linear algebra. The user can visit the Eigen website (https://eigen.tuxfamily.org/dox/), or use their system's package manager. For example on Mac OS with homebrew,

% brew install eigen

# 2.2 Downloading the Repository

The software requires downloading the BAND framework software repository into some directory. Should that be in the User's home directory, the User might enter

/Users/CarlosSmith% git clone https://github.com/bandframework/bandframework.git

Within the repository, there will be a directory

/Users/CarlosSmith/bandframework/software/SmoothEmulator/. All the relevant functionality of *Smooth Emulator* is contained within this directory. Throughout the manual the phrase \${GITHOME\_BAND\_SMOOTH} will refer to this directory.

The User needs to create a personal directory from which the User would perform most projects. This is easiest accomplished by copying a template from the *Smooth* distribution. For example

% cp -r \${GITHOME\_BAND\_SMOOTH}/templates /Users/CarlosSmith/mysmoothy

Within this directory one will find a template project directory, which for the template would then be /Users/CarlosSmith/mysmoothy/myproject. Hence forth, \${MY\_PROJECT} is a name chosen by the User, and will refer to this directory, including the path, from which the User will perform most of the analysis. The User may wish to have several such directories, located according to the User's preference. Any time a new analysis is performed with new parameters, and if the User wishes to save the previous analysis, a new \${MY\_PROJECT} directory should be created. These directories should be outside the main distribution, i.e. outside the bandframework/ path so that the User's work does not interfere with the repository.

Within the templates/ directory, there is also a directory ../templates/mylocal/, which after the copying above will become: /Users/CarlosSmith/mysmoothy/mylocal. The mylocal/ directory contains the main programs and the *Smooth Emulator* executables which will be stored in \${GITHOME\_BAND\_SMOOTH}/mylocal/bin. Because the User might wish to edit the main programs, or to add similar programs, this provides the User a space to make such edits, all while leaving an original copy of the directory in the templates/ directory.

For the remainder of this manual, \${GITHOME\_BAND\_SMOOTH}, \${MY\_LOCAL} and \${MY\_PROJECT} will be used to denote the location of these directories.

# 2.3 Directory and File Structure

The \${GITHOME\_BAND\_SMOOTH}/software/ directory contains codes that are used to create libraries specific to the sampler and emulator. The User can change to The executables are stored in \${MY\_LOCAL}/bin. The short main program source files are located in

\$\{\text{MY\_LOCAL}\}\software\/\main\_\text{programs}. It is not envisioned that the User would edit files in the SmoothEmulator\/\software\/\directory\, but that the User may well wish to create custom versions of the short main programs in \$\{\text{MY\_LOCAL}\}\/\software\/\main\_\text{programs}\/\. The main programs are compiled using the CMake files in \$\{\text{MY\_LOCAL}\}\/\software\/\main\_\text{programs}\/\. The User may find it convenient to add \$\{\text{MY\_LOCAL}\}\/\bin\/\ to their path.

# 2.4 Compiling Libraries

First, change into software directories, then create the makefiles with cmake, then compile them.

```
% cd ${GITHOME_BAND_SMOOTH}/software
${GITHOME_BAND_SMOOTH}/software% cmake .
${GITHOME_BAND_SMOOTH}/software% make
```

There seems to be a common problem that **cmake** misreports the path of the **Eigen** installation. If the User should get an error stating that the Eigen header files cannot be found, the User can set the environmental variable,

```
% export EIGEN3_INCLUDE_DIR=/usr/local/include/eigen3
```

The final arguments may need to be changed depending on the User's location of the packages. If the User wishes to choose a specific C++ compiler, the cmake command should be replaced with cmake -D YOUR\_PREFERRED\_COMPILER.

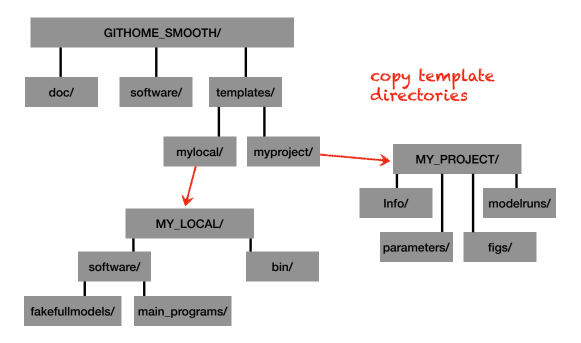


Figure 2.1: The directory structure: The User clones the repository into some location, which will be referred to as \${GITHOME\_BAND\_SMOOTH}. The User can then copy the \${MY\_PROJECT} directory to the User's choices of locations outside the path of the BAND repository. The programs are designed to be run from within the \${MY\_PROJECT}/ directory, which also where data files related to the project will be located. The User can make multiple copies of \${MY\_PROJECT} whenever they wish to run and save different instances of Smooth Emulator. All Smooth Emulator executables are designed to be run from the \${MY\_PROJECT}/ directory. Within that directory there is a subdirectory, \${MY\_PROJECT}/smooth\_data/, within which the data files used by Smooth Emulator reside. The actual executables will be stored in the \${MY\_LOCAL}/bin directory. The main programs are also located in this path, while the libraries will be located in the main repository path. This design was made with the idea that the User might wish to create their own versions of the main programs, and store them outside the repository structure.

At this point all the libraries are built, but this does not include the main programs. The main programs are short, and are meant to serve as examples which the User might copy and edit at will. Before compiling, one needs to set an environmental variable so that the compilation can find the libraries needed for compilation.

% export GITHOME\_BAND\_SMOOTH=/Users/CarlosSmith/bandframework/software/SmoothEmulator

The first part of the path needs to be replaced with the location of the bandframework git repository. If one prefers not to set an environmental variable, one can instead edit \$\{MY\_LOCAL\}/software/CMakeLists.txt, and replace the line where the variable GITHOME\_BAND\_SMOOTH is set to the shell's environmental variable with one where it is set explicitly.

Below, this illustrates how to compile the programs used for generating training points with Simplex and for tuning the emulator with *Smooth Emulator*:

```
% cd ${MY_LOCAL}/software
${MY_LOCAL}/software% cmake .
${MY_LOCAL}/software% make
.
```

The cmake command will also recompile the main libraries in \${GITHOME\_BAND\_SMOOTH}/software/ if necessary. Several source codes for main programs can be found in \${MY\_LOCAL}/software/main\_programs/. If you build your own main programs (probably using these as examples), you can edit the CMakeList.txt file in \${MY\_LOCAL}/software/main\_programs/, using the existing entries as an example. The executables should appear in \${MY\_LOCAL}/bin/.

# 2.5 The MY\_PROJECT Directory

Within \${MY\_PROJECT}/smooth\_data there are several sub-directories (assuming it was created from the template). The first is smooth\_data/Info/. Information about the model parameters, and their priors is stored in smooth\_data/Info/modelpar\_info.txt, and information about the observables is stored in smooth\_data/Info/observable\_info.txt. The file smooth\_data/Info/experimental\_info.txt stores information about the measurements for each observable, and is not needed for building the emulator, but is used by the MCMC investigation of the posterior.

The smooth\_data/smooth\_parameters directory stores user-defined parameter files used by Simplex Sampler, smooth\_data/smooth\_parameters/simplex\_parameters.txt, and by Smooth Emulator smooth\_data/smooth\_parameters/emulator\_parameters.txt. These are user-defined parameters related to choices in running the emulation programs, not model parameters.

The directories smooth\_data/modelruns/run0/, .../modelruns/run1, ..., have files describing the model parameters for each run, along with the output required by the emulator for each specific full-model run. For example, the smooth\_data/modelruns/run1/ directory has the files mod\_parameters.txt and obs.txt. The first file stores the model parameter values for that particular training run. The User then runs their full model based on those parameters and stores the corresponding observables in obs.txt. The User may generate the mod\_parameters.txt files using

 $Simplex\ Sampler$ , or the user might generate them according to some other prescription. Once the User has then generated the obs.txt files,  $Smooth\ Emulator$  can then build and tune the emulator.

Finally the \${MY\_PROJECT}/figs directory contains python scripts for plotting results.

# 3 Generating Training Points with Simplex Sampler

# 3.1 Summary

Simplex Sampler produces a list of points in the n-dimensional model-parameter space to be used for training an emulator. The algorithms are based on the n-dimensional simplex. For example, in two dimensions the points are arranged in an equilateral triangle, and for three dimensions of parameters points are arranged in a tetrahedron. The program first reads a simple text file that provides the names of model parameters and the range of their prior distribution. Options for Simplex sampler are taken from a separate text file. This enables the User to make choices, such as which algorithm to apply when generating the training points. Simplex Sampler determines the number of training points based on the algorithm. The User is free to run the full model at additional points, or to use their own method to generate training points.

# 3.2 Simplex Parameters (not model parameters!)

These are parameters representing choices made by the User. Note these are NOT the model parameters, which are then generated by Simplex Sampler. If one visits the User's project directory, these parameters are stored in the file smooth\_data/smooth\_parameters/simplex\_parameters.txt, where the path is either absolute or relative to the project directory. Parameters files can have any name or location. These files are text files in the format. An example of a parameter file is:

#LogFileName simplexlog.txt # if commented, output to screen
Simplex\_TrainType 1 # Must be 1 or 2
Simplex\_ModelRunDirName modelruns # Directory with training
point information
lea

For the parameter file, the first string is the parameter name and is followed by the value. Both are single strings (without spaces). The # symbol is used for comments. Each parameter has a default value, which will be used if the parameter is not mentioned in the parameter file. Simplex Sampler has four User-defined parameters.

## 1. Simplex\_TrainType

Possible values are "1" or "2". The default, "1", will position points according to a simplex, i.e. in two dimensions this is an equilateral triangle and in three dimensions, it is a tetrahedron. In n dimensions there are n+1 points separated at equal distances from one another and centered at the origin. For "2", points are added at the half-way points between each vertex of the tetrahedron. The points at the bisection points are scaled to a different radius than those at the vertices. This provides the precise number of training points to exactly determine both the linear and quadratic terms.

## 2. Simplex\_ModelRunDirName

This sets the path to the directory in which the run files will be created. The default name is ./modelruns, but the User can change this to anything they want. The path is relative to the project directory, i.e. the directory from which you run the *simplex* command.

## 3. LogFileName

If this is left blank, *Simplex Sampler* will write output to the string. Otherwise it will write output to a file. Given that *Simplex Sampler* runs in a few seconds, the program is usually run interactively and output is sent to the screen.

# 3.3 Specifying Model Parameters and Priors

Before proceeding, Simplex requires information about the parameters, specifically, their ranges. The User enters this information into the file smooth\_data/Info/modelpar\_info.txt. An example of such a file might be

NuclearCompressibility	gaussian	210	40
ScreeningMass	uniform	0.3	1.2
Viscosity	uniform	0.08	0.3

The first column is the model-parameter name, and the last three parameters describe the range of the parameters, which is usually the prior, assuming the prior is uniform or Gausian. The second entry for each parameter defines whether the range/prior is uniform or gaussian. If the prior is uniform, the next two numbers specify the lower and upper ranges of the parameter. If the range/prior is gaussian, the third entry describes the center of the Gaussian,  $x_0$ , and the fourth entry describes the Gaussian width,  $\sigma_0$ , where the prior distribution is  $\propto \exp\{-(x-x_0)^2/2\sigma_0^2\}$ . Simplex will read the information to determine the number of parameters. It will then assign the n points,  $\theta_{1...n}$  assuming each dimension of  $\theta$  varies from -1 to 1, for uniform distributions, or proportional to  $e^{-\theta^2/2}$  for Gaussian distributions. The points  $\theta_i$  are each then converted into  $x_i$  by scaling and translating the values according to the ranges/priors defined in the modelpar\_info.txt file.

# 3.4 Training Types

# 3.4.1 Type 1

Depending on the number of parameters, n, the program creates a simplex in n dimensions. This simplex's vertices will be used to generate  $N_{\text{train}} = n + 1$  training points. These points will be scaled by different values so the training points aren't in the same radius. This results in the minimum number of required points for linear fits. Thus, if the model is perfectly linear, this option provides perfect emulation.

## 3.4.2 Type 2

Depending on the number of parameters, the program first creates a simplex in n dimensions. This simplex's vertices will be used to generate new training points there and along the edges. These points will be scaled to be in different radii from the center. This results in the minimum number of required points for quadratic fits. The net number of training points is then  $N_{\text{train}} = n + 1 + n(n + 1)/2$ . Thus, if the model is perfectly quadratic, this option provides perfect

emulation. Rather arbitrarily, the points generated from the midpoints of the original simplex pairs are all pushed out to a large radius, and those from the original simplex are brought somewhat inward.

# 3.5 Running Simplex to Generate Training Points

To run Simplex Sampler, first make sure the program is compiled. To compile the programs, change into the MY\_LOCAL/main\_programs/ directory and enter the following command,

```
${MY_LOCAL}/software% cmake .
${MY_LOCAL}/software% make simplex
```

Next, change into your project directory and run the program.

```
${MY_PROJECT}% ${MY_LOCAL}/bin/simplex
```

Here \$\{MY\_LOCAL\}/bin is the path to where the User compiles the main programs into executables.

Simplex will read parameters from the smooth\_data/smooth\_parameters/simplex\_parameters.txt file and from the smooth\_data/Info/modelpar\_info.txt files. It will then write the information about the training points in the directory smooth\_data/modelruns/. Within the directory, a subdirectory will be created for each training point, named run0/, run1/, run2/···. Within each subdirectory, Simplex creates a file runI/mod\_parameters.txt for the I<sup>th</sup> training point. For example, the run0/mod\_parameters.txt file might be

NuclearCompressibility	229.08
ScreeningMass	0.453
Viscosity	0.192

At this point, it is up to the User to run their full model at each training point and create a file runI/obs.txt, which stores values of the observables at those training points as calculated by the full model.

# 3.6 Replacing Simplex Sampler with Other Choices of Training Points

If the User desires to use their own procedure for training points, or if the User wishes to augment the Simplex Sampler points with additional training points, the User can write their own files runI/mod\_parameters.txt. The emulator should work fine, though the User should remember that when training the emulator (see Sec. 5) the emulator parameter describing with runI directories to apply should be modified.

There is one warning to be issued when choosing training points for tuning *Smooth Emulator*. When solving for the Taylor coefficients, the emulator uses the factor of those coefficients (products of  $\theta_i$ ) in the linear algebra routine. However, if those coefficients are not linearly independent, the solution becomes undetermined. For example, if two of the training points were exactly the same, it would fail.

# 4 Performing Full Model Runs

There are two forms for storing training data. In the default form, where the parameter SmoothEmulator\_TrainingFormat is set to training\_format\_smooth, training data for a run generated from a specific point in model-parameter space is stored in a unique directory. To be more compatible with the SURMISE statistical package, there is a second option, where SmoothEmulator\_TrainingFormat is set to training\_format\_surmise. For this option the model-parameter values for all the training runs are stored in a single file, defined by the parameter SmoothEmulator\_TrainingThetasFilename and the model observables are stored in a single file SmoothEmulator\_TrainingThetasFilename. These parameters are defined in smooth\_data/smooth\_parameters/emulator\_parameters.txt.

# 4.1 Default Format for Training Data

Once the training points are generated, the User must run the full model for each of the training points. At this point there is a directory, usually called  ${MY_PROJECT}/smooth_data/modelruns/$ , in which there are sub directories, run0/, run1/, run2/.... Within each sub-directory, run1, there should exist a text file smooth\_data/modelruns/run1/mod\_parameters.txt, where  $I = 0, 1, 2, \cdots$ . These files could have been generated by  $Simplex\ Sampler$ , but could have been generated by any other means, including by hand. The files should be of the form,

```
par1_name par1_value
par2_name par2_value
par3_name par3_value
:
```

The parameter names must match those defined in \$\{MY\_PROJECT\}/Info/modelpar\_info.txt, the format of which is described in Sec. 3.

The User must then perform full model runs using the model-parameter values as defined in each sub-directory. The full model runs should record results by writing a list of observable values in each run directory. Each file must be named  $smooth_data/modelruns/runI/obs.txt$ , where  $I = 0, 1, 2, \cdots$ . The format of those text files should be

```
observable1_name observable1_value observable1_random_uncertainty observable2_name observable2_value observable2_random_uncertainty observable3_name observable3_value observable3_random_uncertainty .
```

The names must match those listed in smooth\_data/Info/observable\_info.txt, which will be used by Smooth Emulator, as described in Sec. 5. The values are the observable values as calculated by the full model for the model-parameter values listed in the corresponding mod\_parameters.txt file in the same directory.

The random uncertainties refer only to those uncertainties due to noise in the full model. Random noise is that, which if the full model would be rerun at the same model-parameter values, would

represent the variation in the observable values. In most cases this would be set to zero. But, if the full model has some aspect of sampling to it, for example generating observables from event generators with a finite number of events, that variation should be listed here. This variation is required for the emulator. If there is such a variation, the User might not wish to constrain the emulator to exactly reproduce the training point observables at the training points. The principal danger being, that if two training points are very close to one another, but with a finite fluctuation, exactly producing the training points might require very high slopes to exactly reproduce the training points. If the training points are far apart from one another, and if the random uncertainties are not large, it should be safe to ignore the random uncertainty and constrain the emulator to exactly reproduce the model values.

Once the observable files are produced for each of the full model runs, the User can then proceed to build and tune an emulator using *Smooth Emulator*.

# 4.2 SURMISE Compatible Format for Training Data

In this case the file defined by the parameter SmoothEmulator\_TrainingThetasFilename. For example, this might be set to TrainingThetas.txt, which would then be located in the top level of the project directory. The User would construct the file in the form

```
X1 X2 X3 ....
X1 X2 X3 ....
:
```

where each row listed the model parameters for a specific training run. Similarly, the parameter SmoothEmulator\_TrainingThetasFilename would describe the file where the observables from the training runs were listed. For example, the parameter might be set to TrainingObs.txt. The format might look like

```
Y1 Y2 Y3 ....
Y1 Y2 Y3 ....
```

Again, each row corresponds to a different training point. The two files should thus have the same number of rows.

With this option, there is no way to choose a subset of points from which to train.

# 5 Tuning the Emulator

# 5.1 Summary

Smooth emulator finds an optimum set of Taylor expansion coefficients that reproduce a set of observables at a set of training points. The process of finding those coefficients is referred to as "tuning". For a given observable, a particular sample set of coefficients gives the following emulated function:

$$E(\vec{\theta}) = \sum_{\vec{n}, s.t. \sum_{i} n_{i} \leq \text{MaxRank}} d(\vec{n}) A_{\vec{n}} \left(\frac{\theta_{1}}{\Lambda}\right)^{n_{1}} \left(\frac{\theta_{2}}{\Lambda}\right)^{n_{2}} \cdots .$$
 (5.1)

Here,  $\theta_1\theta_2\cdots$  represent the original model parameters,  $\vec{X}$ , but are scaled. If their initial prior is uniform, they are scaled so that their priors range from -1 to +1, and if they have Gaussian priors, they are scaled so that their variance is one third. The degeneracy factor,  $d(\vec{n})$  is the number of different ways to sum the powers  $n_i$  to a given rank,

$$d(\vec{n}) = \sqrt{\frac{(n_1 + n_2 + \cdots)!}{n_1! n_2! \cdots}}.$$
 (5.2)

As described in Sec. 9, the coefficients are chosen weighted by the distribution,

$$P(\vec{A}) = \prod_{n} \frac{1}{\sqrt{2\pi\sigma_A^2}} e^{-A_n^2/2\sigma_A^2},$$
 (5.3)

where  $\sigma_{\Lambda}$  is varied to maximize the overall probability given the constraint of reproducing the training points. More discussion is provided in Sec. 9. Whereas *Smooth Emulator* does a nice job of finding an optimum value for  $\sigma_{\Lambda}$  if  $\Lambda$  is known, the smoothness parameter  $\Lambda$  is unfortunately difficult to optimize. For the moment, this is treated purely as prior knowledge, or expectation. If the User expects the full model to be very smooth, i.e. the quadratic contributions to be much smaller than the linear contributions and so on, a larger value (e.g. 5.0), might be chosen. If the full-model output might be almost wavy, then a smaller value (e.g. 2) might be chosen. The emulator uncertainties will be smaller for larger  $\Lambda$ .

By setting parameters, as described below, *Smooth Emulator* can be tuned one of three different ways

- a) Find the optimum set of coefficients. If evaluated at the training points, the emulator will exactly produce the full model. When it predicts the observable at a new  $\vec{\theta}$  it provides an uncertainty.
- b) If a Monte Carlo tuning method is chosen, the emulator finds a predetermined number of sets of coefficients, where each set of coefficients provides a function that exactly reproduces the full model at the training points. Aside from the constraint, the coefficients are chose randomly, but weighted according to Eq. (5.3). The User sets the number of sets of coefficients, typically of order  $N_{\text{sample}} \approx 10$ , in the parameter file. Away from the training points, the uncertainty of the emulator is represented by the spread of the values amongst the  $N_{\text{sample}}$  predictions.

c) The third mode also provides  $N_{\text{sample}}$  predictions, but rather than exactly reproducing the training values the emulator merely comes close to the training points with a distribution  $\sim e^{-\Delta y^2/2\epsilon_y^2}$ , where  $\epsilon_y$  represents the random error of the full model. This mode should be chosen if the full model has significant random error, and especially if the training points are close to one another.

Method (a) is by far the quickest, and will probably be used the most often.

If methods (b) or (c) are chosen  $Smooth\ Emulator$  solves for the  $N_{sample}$  sets of coefficients from the training data, then stores  $N_{sample}$  sets of coefficients, along with the averaged coefficients in files for later use. If (a) is chosen,  $Smooth\ Emulator$  stores the set of "best" coefficients along with some other arrays used for rapid calculation of the uncertainty.  $Smooth\ Emulator$  can emulate either the full-model observables directly, or their principal components. Training the emulator follows the same steps for either approach.

The executables based on *Smooth Emulator* are located in the User's \${MY\_LOCAL}/bin directory. Examples of such executables are smoothy\_tune or smoothy\_calcobs. These functions must be executed from within the User's project directory.

In the following subsections, we first review the format for each of the required input files, then describe how to run *Smooth Emulator*, how its output is stored, and how to switch PCA observables for real observables.

# **5.2** Preparing Files for Smooth Emulator

Before training the emulator, one must first run the full model at a given set of training points. In addition to a parameter file (described in the next sub-section), which sets numerous options, the User must provide the following:

 A file listing the names of observables. This file is named smooth\_data/Info/observable\_info.txt, where the path is relative to the project directory. The file might look like

```
obsname1
obsname2
obsname3
obsname4
:
```

2. A file listing the names of the model parameters that also describes their priors. This file is smooth\_data/Info/modelpar\_info.txt. The file might have the following form:

```
parname1 uniform 0 1.0E-3
parname2 uniform -50.0 100.0
parname3 gaussian 0 24.6
parname4 uniform 30.0 50.0
```

:

If the prior is uniform the two following numbers provide the minimum and maximum of the interval. If the prior is gaussian the two subsequent values represent the center and r.m.s. width of the Gaussian. This same file was required for running *Simplex Sampler*.

3. The User must provide a list of the model-parameter values,  $\vec{\theta}_{train}$ , at each training point. These points can be generated by  $Simplex\ Sampler$ , as described in Sec. 3, or they can be generated by hand.

There are two formats for entering the information, and the User chooses between them by changing the SmoothEmulator\_TrainingFormat parameter described below. The standard format involves writing the training output in directories, run1,run2,... with the smooth\_data/modelruns/ directory. If the number of full-model runs performed is  $N_{\text{train}}$ , Smooth emulator requires files for each run. Each file is named smooth\_data/modelruns/run1/mod\_parameter  $0 \le I < N_{\text{train}}$ , and I denotes the point in parameter space for the  $I^{\text{th}}$  full-model training run. mod\_parameters.txt might look like

```
parname1 8.34E-4
parname2 -30.5375
parname3 36.238
parname4 39.34
:
```

The alternative format involves writing the model output in a single file specified by the SmoothEmulator\_ThetasFileName parameter. That format is

```
X1 X2 X3 ....
X1 X2 X3 ....
:
```

The number of rows corresponds to the number of training points. There is no option to choose a subset of the training points with this option. This format is chosen by setting SmoothEmulator\_TrainingFormat to training\_format\_surmise.

4. At each training point, the User must provide the full model's value for each observable. Again, there are two options, with the choice defined by the SmoothEmulator\_TrainingFormat parameter. For the default format:

In the same directory where the model-parameter values are stored, *Smooth Emulator* requires the observables calculated at the training points mentioned above. This information is provided in <code>smooth\_data/modelruns/runI/obs.txt</code>. An example of such a file is:

```
obsname1 -51.4645 2.5
obsname2 166.837 0.9
obsname3 -47.9877 0.0
obsname4 -2.34526 0.03
```

:

The first number is the calculated value of the observable, and the second is the random error. This is only the random error, i.e. that which represents that if the model were rerun at the same training point, the value might be different. This should only be non-zero if the full-model has some Monte Carlo feature. For example, the full model might involve simulating a small number of events. Other types of uncertainty are accounted for by including them into the experimental uncertainty.

The alternative format is again chosen by setting SmoothEmulator\_TrainingFormat to training\_format In that case a file, specified by the SmoothEmulator\_ObsFileName parameter is of the form:

```
Y1 Y2 Y3 ....
Y1 Y2 Y3 ....
```

# 5.3 Experimental Measurement Information

Once the emulator is tuned and before it is applied to a Markov Chain investigation of the likelihood, the software needs to know the experimental measurement and uncertainty. That information must be entered in the smooth\_data/Info/experimental\_info.txt file. The file should have the format:

```
-12.93
                     0.95
                             0.5
obsname1
          159.3
                     3.0
                             2.4
obsname2
          -61.2.
                     1.52
                             0.9
obsname3
obsname4
          -1.875
                     0.075
                             0.03
```

The first number is the measured value and the second is the experimentally reported uncertainty. The third number is the uncertainty inherent to the theory, due to missing physics. For example, even if a model has all the parameters set to the exact value, e.g. some parameter of the standard value, the full-model can't be expected to exactly reproduce a correct measurement given that some physics is likely missing from the full model. For the MCMC software, the relevant uncertainty incorporates both, and only the combination of both, added in quadrature, affects the outcome. We emphasize that this last file is not needed to train and tune the emulator. It is needed once one performs the MCMC search of parameter space.

# 5.4 Smooth Emulator Parameters (not model parameters!)

Smooth Emulator requires a parameter file,

\${MY\_PROJECT}/smooth\_data/smooth\_parameters/emulator\_parameters.txt. The parameter file is simply a list, of parameter names followed by values.

```
#LogFileName smoothlog.txt # comment out for interactive running
SmoothEmulator_LAMBDA 2.5 # Smoothness parameter
SmoothEmulator_MAXRANK 5
SmoothEmulator_ConstrainAO true
SmoothEmulator_TrainingPts 0-27
SmoothEmulator_TestPts 28-50
SmoothEmulator_UsePCA false
SmoothEmulator_TrainingFormat training_format_smooth
#SmoothEmulator_TrainingFormat training_format_surmise
SmoothEmulator_TrainingThetasFilename TrainingThetas.txt
SmoothEmulator_TrainingObsFilename TrainingObs.txt
#
```

The last two filenames are only relevant if the SURMISE training format is used. If any of these parameters are missing from the parameters file, *Smooth Emulator* will assign a default value.

## • LogFileName

If this is commented out, as it is in the example above, *Smooth Emulator*'s main output will be directed to the screen and the program will run interactively. Otherwise, the output will be recorded in the specified file. Most often, one will wish the program to run interactively.

#### • SmoothEmulator\_LAMBDA

This is the smoothness parameter  $\Lambda$ . It sets the relative importance of terms of various rank. If  $\Lambda$  is unity or less, it suggests that the Taylor expansion converges slowly. The default is 3.

## • SmoothEmulator\_MAXRANK

As Smooth Emulator assumes a Taylor expansion, this the maximum power of  $\theta^n$  that is considered. Higher values require more coefficients, which in turn, slows down the tuning process. The default is 4.

#### • SmoothEmulator\_ConstrainA0

The coefficients in the Taylor expansion are assumed to have some weight,

$$W(A_i) = rac{1}{\sqrt{2\pi\sigma_A^2}}e^{-A_i^2/2\sigma_A^2}.$$

The term  $\sigma_A$  is allowed to vary during the tuning to maximize the likelihood of the expansion. If the User wishes to exempt the lowest term, i.e. the constant term in the Taylor expansion from the weight, the User may set SmoothEmulator\_ConstrainAO to false. The default is false.

### • SmoothEmulator\_TrainingPts

This lists which full-model training runs SmoothEmulator will use to train the emulator. This provides the User with the flexibility to use some subset for training, as may be the case when testing the accuracy. The default is "1". An example the User might enter could be SmoothEmulator\_TrainingPts 0-4,13,15

This would choose the training information from the directories run0, run1, run2, run3, run4, run13 and run15, which would be found in smooth\_data/modelruns/.

#### • SmoothEmulator\_TestPts

This lists which full-model training runs SmoothEmulator will use to test the emulator away from where the emulator was tuned. This provides the User with the flexibility to use some subset for training, as may be the case when testing the accuracy. The default is "1". An example the User might enter could be

```
SmoothEmulator_TestPts 5-12,14,16-50
```

This would choose the model output information from the specified directories, run5, run6... found in

smooth\_data/modelruns/.

## • SmoothEmulator\_TrainingFormat

This defines the format to be used for reading the training data. There are two choices. The default is:

SmoothEmulator\_TrainingFormat training\_format\_smooth

This reads model output from the smooth\_data/modelruns/ directory. The second choice is: SmoothEmulator\_TrainingFormat training\_format\_surmise

This latter choice reads the training information from files defined by the parameters SmoothEmulator\_TrainingThetasFilename and SmoothEmulator\_TrainingObsFilename. For example, the following lines would set the format and read the input from the specified files:

```
SmoothEmulator_TrainingFormat training_format_surmise
SmoothEmulator_TrainingThetasFilename TrainingThetas.txt
SmoothEmulator_TrainingObsFilename TrainingObs.txt
```

The format for the file specified by SmoothEmulator\_TrainingThetasFilename should be:

```
X1 X2 X3 ....
X1 X2 X3 ....
```

Here, each row represents the model parameters for a different training point. All the lines will be used to tune the emulator. The format for the file SmoothEmulator\_TrainingObsFilename should be:

```
Y1 Y2 Y3 ....
Y1 Y2 Y3 ....
```

Again, each row should correspond to a different training point. The two files above should thus have the same number of rows.

#### • SmoothEmulator\_UsePCA

By default, this is set to false. If one wishes to emulate the PCA observables, i.e. those that are linear combinations of the real observables, this should be set to true. One must then be sure to have run the pca decomposition programs first. For more, see Sec. 6.

# 5.5 Running Smooth Emulator Programs

The source code for several *Smooth Emulator* main programs can be found in the \${MY\_LOCAL}/main\_programs/ directory. They are separated from the bulk of the software, which is in the \${bandframework}/SmoothEmulator/software/ directory. The main programs are designed

so that the User can easily copy and edit them to create versions that might be more appropriate to the User's specific needs. When compiled, from the \${MY\_LOCAL}/main\_programs/ directory, the executables appear in the \${MY\_LOCAL}/bin/ directory.

To begin, we consider the source code \${MY\_LOCAL}main\_programs/smoothy\_tune\_main.cc. Once compiled the corresponding executable is \${MY\_LOCAL}/bin/smoothy\_tune. The source code for smoothy\_tune is:

```
int main(){
    // First create CSmoothMaster
    CparameterMap *parmap=new CparameterMap();
    CSmoothMaster master(parmap);
    // Tune for all observables "Y"
    master.TuneAllY();
    // Write coefficients to file
    master.WriteCoefficientsAllY();
    return 0;
}
```

From within the \${MY\_LOCAL}/main\_programs/ directory, one can compile the program with the command:

```
MY_LOCAL/main\_programs % cmake .
MY_LOCAL/main\_programs % make smoothy_tune
```

The executable smoothy\_tune should now appear in the \${MY\_LOCAL}/bin/ directory. If one enters make without the name of the program, all the main-program source files will be compiled. Assuming the directory \${MY\_LOCAL}/bin/ has been added to the User's path, the User may switch to the User's project directory, and enter:

```
~/MY_PROJECT % smoothy_tune
```

This should read in all the necessary information, tune the emulators for all the observables and write the Taylor-expansion coefficients to file. The optimum coefficients are stored in the file smooth\_data/coefficients/OBS\_NAME/ABest.bin, where OBS\_NAME is the observable name. If a Monte Carlo tuning method was applied, there are several sets of coefficients stored, smooth\_data/coefficients/OBS\_NAME/sampleI.bin, where  $0 \le I < N_{\text{sample}}$ . Along with the coefficients, in the same directory  $Smooth\ Emulator$  writes a file for each observable. These files are named smooth\_data/coefficients/OBS\_NAME/meta.txt. This file provides information, such as the maximum rank and net number of model parameters, to make it possible to read the coefficients later on. For the non-Monte-Carlo method, files smooth\_data/coefficients/OBS\_NAME/BetaBest.bin are stored for later use in calculating the uncertainties.

Similarly, there is a code \${MY\_LOCAL}/main\_programs/smoothy\_calcobs\_main.cc, which provides an example of how one might read the coefficients and generate predictions for the emulator at specified points in parameter space:

```
int main(){
  // Create Smooth Master Object
 NMSUUtils::CparameterMap *parmap=new CparameterMap();
 NBandSmooth::CSmoothMaster master(parmap);
  // Read in coefficients
 master.ReadCoefficientsAllY();
  // Create a CModelParameters object to store information
    // about a single point in parameter space
 NBandSmooth::CModelParameters *modpars=new NBandSmooth::CModelParameters();
     // contains info about single point
 modpars->priorinfo=master.priorinfo;
  master.priorinfo->PrintInfo(); // print info about priors
  // Prompt user for model parameter values
  vector<double> X(modpars->NModelPars);
  for(unsigned int ipar=0;ipar<modpars->NModelPars;ipar++){
    cout << "Enter value for " << master.priorinfo->GetName(ipar) << ":\n";</pre>
    cin >> X[ipar];
 modpars->SetX(X);
  // Calc Observables and print out their values and uncertainties
  NBandSmooth::CObservableInfo *obsinfo=master.observableinfo;
  vector<double> Y(obsinfo->NObservables);
  vector<double> SigmaY(obsinfo->NObservables);
 master.CalcAllY(modpars,Y,SigmaY);
  cout << "--- EMULATED OBSERVABLES ----\n";</pre>
  for(unsigned int iY=0;iY<obsinfo->NObservables;iY++){
    cout << obsinfo->GetName(iY) << " = " << Y[iY] << " +/- " << SigmaY[iY] << endl;</pre>
 }
 return 0;
```

Smooth Emulator programs will often output lines describing their progress, either to the screen or to a file, as specified by the SmoothEmulator\_LogFile parameter described above. For example, with the Monte Carlo tuning methods the output includes a report on the percentage of Metropolis steps in the MCMC program that were successful. The line BestLogP/Ndof describes the weight used to evaluate the likelihood of a coefficients sample. This value should roughly plateau once the Metropolis procedure has settled on the most likely region. All output, except for some explicit code in main programs, is directed in this manner.

# 5.6 Useful Functionalities of CSmoothMaster Object

Smooth Emulator was designed so that the User can write their own main programs and access the functionality mainly through references to the CSmoothMaster object. Additionally, the User might find it useful to access the CModelParameters, CparameterMap, CPriorInfo, and CObservableInfo objects. Here is a compendium of calls to the CSmoothMaster:

# • CSmoothMaster(CparameterMap \*parmap)

This is the constructor. CparameterMap object stores temulator parameters (not model parameters). CparameterMap functionality described below.

## • void ReadTrainingInfo()

This reads the training point information to be used for tuning. It is done automatically when creating the CSmoothMaster object.

## • void TuneAllY()

// Tune all observables, Y.

## • void TuneY(string obsname)

Tunes one observable, by observable name.

## • void TuneY(unsigned int iY)

Tune one observable, referenced by index.

• void CalcAllY(CModelParameters \*modelpars, vector <double > &Y, vector <double > &SigmaY\_emulator)

Calculates all observables. CModelParameters object stores information about a single point in model-parameter space. Object described further below.

 void CalcY(unsigned int iY,CModelParameters \*modelpars,double &Y, double &SigmaY\_emulator)

Calculates observable referenced by index.

void CalcY(string obsname, CModelParameters \*modelpars, double &Y, double &SigmaY\_emulator)

Calculates observable referenced by observable name.

- void CalcAllYdYdTheta(CModelParameters \*modelpars, vector<double> &Y, vector<double> &SigmaY\_emulator, vector<vector<double>> &dYdTheta)
   Also calculates derivatives w.r.t. θ. Especially useful for some Markov chain searches in parameter space, e.g. Langevin approaches.
- void CalcYdYdTheta(string obsname, CModelParameters \*modelpars, double &Y, double &SigmaY\_emulator, vector <double > &dYdTheta)
  Same, but for one observable referenced by observable name.
- void CalcYdYdTheta(unsigned int iY, CModelParameters \*modelpars, double &Y, double &SigmaY\_emulator, vector <double > &dYdTheta)
  Same, but by index.
- void CalcAllYOnly(CModelParameters \*modelpars, vector <double> &Y) and CalcAllYOnly(vecto &theta, vector <double> &Y) calculates the observable but with the uncertainty.
- double GetUncertainty(string obsname, vector < double > & theta) and GetUncertainty (unsigned int iY, vector < double > & theta) returns the emulator uncertainty only, for a specific observable.

### • void CalcAllLogP()

Prints technical information the User may find helpful in evaluating whether the choice of  $\Lambda$  is reasonable. For each observable, it calculates the ratio of the r.m.s. coefficients of rank-two to those of rank-one. One would roughly expect the ratio to be unity if  $\Lambda$  is appropriate, but

from testing the measure is so noisy that it is not useful on a observable-by-observable basis. It also calculates the probability for the optimum coefficient set. This would be maximized for best choices of  $\Lambda$ , but again is highly sensitive to fluctuations. Thus this information is not recommended for actual tuning  $\Lambda$ .

## • void TestAtTrainingPts()

Compares emulator predictions to full model calculations at training points. Observables should match and uncertainties should vanish.

# • void TestAtTrainingPts(string obsname)

Same but for a single observable referenced by observable name.

# • void TestAtTrainingPts(unsigned int iY)

Same but for a single observable referenced by index.

#### • void TestVsFullModel()

This is useful for comparing emulator to model evaluated at points not used for training. You can set the emulator parameter SmoothEmulator\_TestPts in the same manner as was done for the emulator parameter SmoothEmulator\_TrainingPts. It will compare the emulator to the results in the run directories specified here. Usually, one wishes to compare at points not used for training. Fore example, if there were 100 run directories, one might set

```
SmoothEmulator_TrainingPts 0-89
SmoothEmulator_TestPts 90-99
```

to train the emulator with the first 90 points and test it at the last 10 points. Tests for each observable are stored in a directory fullmodel\_testdata/.

## • void TestVsFullModelAlt()

This is useful for comparing emulator to model evaluated at points not used for training. For the full model, make a sub-directory within the project directory called fullmodeltestdata/. Within that subdirectory, for each observable create a file called OBSNAME.txt, where OBSNAME is the name for each observable. Each file should have the format

```
X_1 X_2 ... X_N Y
X_1 X_2 ... X_N Y
X_1 X_2 ... X_N Y
```

Each line describes a point in parameter space and the observable calculated by the full model. master.TestVsFullModel() will calculate the emulated value at each  $\vec{X}$  and compare the full-model value to the emulator value. It will also give the uncertainty. If the uncertainty is well represented, 68 % of the emulated values should be within the uncertainty.

#### • void WriteCoefficientsAllY()

Writes the Taylor coefficients for all observables. Because tuning is so fast, the User can usually avoid reading and writing coefficients, and instead simply re-perform the tuning.

## • void WriteCoefficients(string obsname)

Writes only those for one observable referenced by observable name.

- void WriteCoefficients(unsigned int iY)
  Writes only for one observable reference by index.
- void ReadCoefficientsAllY() Reads all Taylor coefficients from file.
- void ReadCoefficients(string obsname)
  Reads Taylor coefficients for one observable referenced by observable name.
- void ReadCoefficients(unsigned int iY)
  Reads Taylor coefficients for one observable referenced by observable index.

It would probably be useful to view the header file for the master class, \$GITHOME\_BAND\_SMOOTH/software/incl

# 5.7 Other Potentially Useful Smooth Emulator Objects

Within the main program one may also wish to apply or access other objects with the *Smooth Emulator* framework. Their functionalities are described here:

## • CparameterMap

This object stores the emulator parameters described above. It is a simple inheritization of a map, linking string labels to values of various types. The object can read a parameter list from a file, e.g.

```
CparameterMap parmap;
parmap.ReadParsFromFile("smooth_data/smooth_parameters/emulator_parameters.txt")
```

The argument can be either a C++ string or a character array. The CSmoothMaster constructor takes a pointer to a CparameterMap object as a argument. If one wishes to print the parameters, the function is parmap.PrintPars(). To set a parameter within a program, parmap.set(string,value), where value can be any of several types, e.g. bool, double, an integer, long long integer, string, .... To retrieve a value from the map, the commands are getB, getI, getLongI, getS, getD, ..., for bool, int, long long int ... types. For example,

```
parmap.getD("SmoothEmulator_LAMBDA",2.0);
```

would retrieve the value of  $\Lambda$ , and if the map did not include SmoothEmulator\_LAMBDA, it would return a default value of 2.0.

#### • CPriorInfo

This object stores information about the prior. The CSmoothMaster object includes such an object, and automatically, during its construction, the CSmoothMaster object reads in the smooth\_data/Info/modelpar\_info.txt file and creates the object. The functionalities of potential interest might be addressed from a main program via:

```
smoothmaster.priorinfo.PrintInfo(); // prints out the parameter priors
unsigned int ipar=smoothmaster.priorinfo.GetIPosition(PARAMETER_NAME);
   // finds position given name of parameter (string)
string parname=smoothmaster.priorinfo.GetName(I);
   // finds name given position {\tt I} (unsigned int)
```

#### • CModelParameters

This object stores the information describing a single point in the model-parameter space. It has two vectors storing the true  $\vec{X}$  and the scaled  $\vec{\theta}$  parameters. As a static variable, it stores a pointer to a CPriorInfo object so that it can translate back and forth from  $\vec{X}$  to  $\vec{\theta}$ . The functionalities are fairly easily seen from the definition of its members header file. The ones most likely to be accessed by the User are:

```
vector<double> X;  // model parameters
vector<double> Theta; // scaled model parameters
void TranslateTheta_to_X();  // Given Theta, fill out X
void TranslateX_to_Theta();  // Given X, fill out Theta
void Print();  // Prints model parameters
void Write(string filename);  // Writes model parameters
void Copy(CModelParameters *mp);  // Copies from another object
void SetTheta(vector<double> &theta);  // Sets model parameters from vector, also translates
```

The emulator software stores all model-parameter information in these objects. There is a CTrainingInfo object within CSmoothMaster that stores a vector of

## • CObservableInfo

This object stores general information about all the observables, including their experimental value and experimental uncertainty. The object does not store observable information as calculated by the emulator. These objects are also fairly self-explanatory and the functionality can be ascertained by looking at some of the lines in the header file.

```
unsigned int NObservables;
vector<string> observable_name;
vector<double> SigmaA0;
  // initial guess for spread of coefficients (only used for MC tuning methods)
map<string,unsigned int> name_map;
unsigned int GetIPosition(string obsname);
  // finds position given name of observable
string GetName(unsigned int iposition);
  // finds name give position
void ReadObservableInfo(string filename);
  // reads information about observable, either from
  //smooth\_data/Info/observable_info.txt or smooth\_data/Info/pca_info.txt
void ReadExperimentalInfo(string filename);
  // reads experimental measurement and uncertainty (used in MCMC)
vector<double> YExp,SigmaExp;
```

```
// experimental measurement information
void PrintInfo();
```

There is such an object in the CSmoothMaster class. For example, to print the information about the observables from a main program, one would include the line: master->observableinfo->PrintInfo();

# 6 Emulating Principal Components

User Beware: the PCA software is under development and is far from being fully tested. Explore at your own risk.

## 6.1 Overview

Rather than emulating all observables, it can be more efficient to emulate a handful of principal components. After generating the training-point data, one can run the pca\_calctransformation program included with the distribution. This will create files that shadow those used to emulate the observables. This will create a file PCA\_Info/observable\_info.txt which shadows the Info/observable\_info.txt. The difference is that the observables will be named z1,z2.... In each run directory, alongside the obs.txt files, there will be a obs\_pca.txt file. Finally, there will be a file PCA\_Info/tranformation\_info.txt file that contains all the information and matrices required to perform the basis transformation. If the parameter Use\_PCA is set to true, the emulator will use the PCA files above instead of the observable files. The emulator will then store the Taylor coefficients in the directory coefficients\_pca/ rather than in coefficients/.

# 6.2 Compiling and Running the PCA Programs

To get an idea of the capabilities and functionality of the PCA elements of the *Smooth Emulator* Distribution, one can view the sample main program,

\${MY\_LOCAL}/main\_programs/pca\_calctransformation\_main.cc:

```
int main() {
   CparameterMap parmap;
   NBandSmooth::PCA *pca = new NBandSmooth::PCA(&parmap);
   pca->CalcTransformationInfo();
}
```

To compile the program,

\${MY\_LOCAL}/main\_programs% make pca\_calctransformation

Before running one needs to have set up the usual files defining the training points, e.g. running Simplex Sampler, and running the full model at the training points. One can now run the program that creates files containing all the observable information, but translated into PCA components:

```
${MY_PROJECT}% ${MY_LOCAL}/bin/pca_calctransformation
```

One can check the directory \${MY\_PROJECT}/PCA\_Info/ to make sure that one sees the files experimental\_info.txt, observable\_info.txt and transformation\_info.txt. The latter file includes the transformation information so that one can readily translate from the nominal observables to the PCA components.

Before performing the tuning, one needs to edit the parameters/emulator\_parameters.txt file and change the parameter SmoothEmulator\_UsePCA to true. The output when tuning the emulator should look something like:

```
${MY_PROJECT}% ${MY_LOCAL}/bin/smoothy_tune
---- Tuning for z0 ----
---- Tuning for z1 ----
---- Tuning for z2 ----
---- Tuning for z3 ----
---- Tuning for z4 ----
:
```

The Taylor-expansion coefficients for the PCA observables are in the directory \${MY\_PROJECT}/coefficients\_p Another example program for PCA analysis is \${MY\_LOCAL}/main\_programs/pca\_readinfo\_calcy\_main.cc:

```
int main() {
CparameterMap parmap;
   NBandSmooth::PCA *pca = new NBandSmooth::PCA(&parmap);
   pca->ReadTransformationInfo();
vector<double> Z,Y,SigmaZ_emulator,SigmaY_emulator;
Z.resize(pca->Nobs);
Y.resize(pca->Nobs);
SigmaZ_emulator.resize(pca->Nobs);
SigmaY_emulator.resize(pca->Nobs);
printf("---- Start with these values of Z ----\n");
for(unsigned int iobs=0;iobs<pca->Nobs;iobs++){
 Z[iobs]=iobs;
printf("Z[%u]=%g\n",iobs,Z[iobs]);
}
printf("--- Translated values of Y ----\n");
SigmaZ_emulator[0]=SigmaZ_emulator[1]=SigmaZ_emulator[2]=SigmaZ_emulator[3]=SigmaZ_emulator
pca->TransformZtoY(Z,SigmaZ_emulator,Y,SigmaY_emulator);
for(unsigned int iobs=0;iobs<pca->Nobs;iobs++){
printf("%10.5f %10.5f\n",Y[iobs],SigmaY_emulator[iobs]);
printf("--- (Re)Translated values of Z ----\n");
pca->TransformYtoZ(Y,SigmaY_emulator,Z,SigmaZ_emulator);
for(unsigned int iobs=0;iobs<pca->Nobs;iobs++){
printf("%10.5f %10.5f\n",Z[iobs],SigmaZ_emulator[iobs]);
printf("--- Retranslated values of Z should match original and SigmaZ should all be unity
}
```

This program illustrates how one can insert the PCA transformations into another program. After defining the pca object, the program reads in the transformation information with the pca->ReadTransformation.

command. The remainder of the program simply test the transformation by making a fake vector of the Z (PCA) components, then translating them to Y (observable) values and then back. The original values for Z[i] are set to  $0, 1, 2, \cdots$ , which should then match the values after translating back and forth as illustrated.

To compile the program:

```
${MY_LOCAL}/main_programs% make pca_readinfo_calcy
```

Running the program:

```
${MY_PROJECT}% ${MY_LOCAL}/bin/pca_readinfo_calcy
---- Start with these values of Z ----
Z[0]=0
Z[1]=1
Z[2]=2
Z[3]=3
Z[4]=4
Z[5]=5
---- Translated values of Y and SigmaY ----
  -2.09833
             24.99999
  -1.03694
             33.33329
  -0.26087
             41.66668
  -5.36371
              0.16667
  3.49439
              0.08333
              0.08333
   2.91093
---- (Re)Translated values of Z and SigmaZ ----
  0.00000
              1.00000
   1.00000
              1.00000
   2.00000
              1.00000
   3.00000
              1.00000
   4.00000
              1.00000
              1.00000
   5.00000
---- Retranslated values of Z should match original and SigmaZ should all be unity ----
```

Note that for the PCA observables the scaling ensures they all have the same uncertainty, 1.0.

# 6.3 PCA Parameters (not model parameters!)

The PCA programs uses parameter that are prefixed with **SmoothEmulator**. One would typically use the same parameter file as used for running *Smooth Emulator*. The relevant parameters are:

#### 1. SmoothEmultor\_UsePCA

If one wishes to emulate the PCA observables, i.e. those that are linear combinations of the real observables, this should be set to true. One must then be sure to have run the PCA decomposition programs first.

2. SmoothEmulator\_ModelRunDirName and SmoothEmulator\_TrainingPts should be set the same as used by Smooth Emulator.

### 3. SmoothEmulator\_PCAMinVariance

This tells the emulator to not emulate PCA observables where the corresponding eigen value to the  $\langle \delta Y_a/\sigma_a \delta Y_b/\sigma_b \rangle$  matrix are below this amount. For such variables the emulated value of  $Z_a$  is simply set to zero.

# 6.4 Performing MCMC search with PCA components

Once the parameter SmoothEmultor\_UsePCA is set to true and the parameter SmoothEmulator\_PCAMinVariance is set to be greater than zero, the MCMC procedure should work in the MCMC programs. Both these parameters are set in the Info/emulator\_parameters.txt file. For example, if SmoothEmulator\_PCAMinVariance is set to 0.01, then the emulator will not bother emulating any PCA observable where, in the training set,  $\langle \delta Z_a \delta Z_a \rangle$  is less than 0.01. Given that the PCA variables are scaled by their uncertainties, this then ignores PCA observables that vary less than one percent of their uncertainty.

The User should be warned again that all this functionality is not yet fully tested.

# 7 Markov-Chain Monte Carlo (MCMC) Generation of the Posterior

# 7.1 Preparing for MCMC using Smooth Emulator

The final step in Bayesian analyses is to generate MCMC traces through parameter space. Smooth Emulator software is designed for the MCMC programs to be run from within the \${MY\_PROJECT} directory. Once the emulator is tuned and before it is applied to a Markov Chain investigation of the likelihood, the software needs know the experimental measurement and uncertainty. That information must be entered in the smooth\_data/Info/experimental\_info.txt file. The file should have the format:

```
-12.93
obsname1
                     0.95
                             0.5
          159.3
                     3.0
                             2.4
obsname2
obsname3
          -61.2.
                     1.52
                             0.9
          -1.875
                             0.03
obsname4
                     0.075
```

The first number is the measured value and the second is the experimentally reported uncertainty. The third number is the uncertainty inherent to the theory, due to missing physics. For example, even if a model has all the parameters set to the exact value, e.g. some parameter of the standard value, the full-model can't be expected to exactly reproduce a correct measurement given that some physics is likely missing from the full model. For the MCMC software, the relevant uncertainty incorporates both, and only the combination of both, added in quadrature, affects the outcome. We emphasize that this last file is not needed to train and tune the emulator. It is needed once one performs the MCMC search of parameter space.

The log-likelihood,  $\boldsymbol{LL}$ , for the MCMC generation is assumed to be of a simple form. Summing over the observables  $\boldsymbol{I}$ ,

$$egin{array}{lll} \sigma_{I,{
m tot}}^2 &=& \sigma_{I,{
m exp}}^2 + \sigma_{I,{
m theory}}^2 + \sigma_{I,{
m emulator}}^2, \ LL &=& -\sum_I rac{(Y_{I,{
m exp}} - Y_{I,{
m emu}})^2}{2\sigma_{I,{
m tot}}^2} - \ln(\sigma_{I,{
m tot}}). \end{array}$$

The main program that runs the mcmc code is \${MY\_LOCAL}/software/main\_programs/mcmc\_main.cc. To compile the program:

### \${MY\_LOCAL}/software% make mcmc

One should tune the emulator before running mcmc. This could involve running the emulator and saving the Taylor coefficients, or inserting the tuning into the main source code for mcmc mentioned above.

## 7.2 MCMC Parameter File

Next, one needs to edit the parameter file \${MY\_PROJECT}/smooth\_data/smooth\_parameters/mcmc\_parameter An example file is:

```
#LogFileName mcmc_log.txt
MCMC_LANGEVIN false
MCMC_METROPOLIS_STEPSIZE 0.04
MCMC_LANGEVIN_STEPSIZE 0.5
MCMC_NBURN 100000
MCMC_NTRACE 100000
MCMC_NSKIP 5
MCMC_IGNORE_EMULATOR_ERROR false
RANDY_SEED 12345
```

As was the case with *Smooth Emulator*, each parameter has a default value. Summarizing the parameters:

#### • LogFileName

To run interactively, leave this line commented out. Otherwise, output will be directed to the designated file.

#### • MCMC\_LANGEVIN

The Langevin method can replace the Metropolis method, but it is not fully tested. If the default, false, is set, the Metropolis method will be invoked. The Langevin method currently ignores the emulator uncertainty. At this time it is not clear that once it is invoked, it will significantly improve performance.

#### • MCMC\_METROPOLIS\_STEPSIZE

Metropolis algorithms require taking random steps in  $\vec{\theta}$  space. If the steps are small, it takes longer to explore the space, but if the steps are very long, the success rate of the Metropolis steps becomes low. Maximum efficiency occurs when the Metropolis success rate, which is provided during running, is near 50%. Rates of 20% or 80% are also fine, but if the rate is only a few percent of if it becomes close to 100%, the User should adjust the parameter.

### • MCMC\_LANGEVIN\_STEPSIZE

For the Langevin procedure, all steps are successful, but the accuracy can suffer if the steps are too large. (The Langevin option is still in development)

#### • MCMC\_NBURN

A certain number of Metropolis or Langevin steps should be taken before the trace is recorded so that the trace is not biased by the starting value.

## MCMC\_NTRACE

This is the number of points in the trace that are recorded for subsequent analysis. More points provides a more accurate representation of the posterior.

#### • MCMC\_NSKIP

Because succesive points in the trace are correlated, it makes sense to skip several points before skipping. For example, in the Metropolis procedure if the success rate is 50%, the neighboring points have a 50% chance of being the same. The values of MCMC\_NBURN, MCMC\_NTRACE and MCMC\_NSKIP are stored by the CMCMC objects, but are only set when the object calls the trace function. Thus, default values are set at the time the call is made to the PerformTrace funtion. See Sec. 7.4 below for an example of how this is used.

#### • MCMC\_IGNORE\_EMULATOR\_ERROR

If this flag is set to true the emulator error will be ignored. This significantly increases the speed of the MCMC procedure, but should not be done if the emulator error is significant.

# 7.3 Running the MCMC Program

To run the provided MCMC program, to the project directory, and run the program mcmc. Output should look something like this:

```
${MY_PROJECT}% ${MY_LOCAL}/bin/mcmc
At beginning of Trace, LL=-68.764478
At end of trace, best LL=1.563806
Best Theta=
0.249554 0.153237 0.190531 0.210907 0.058929 0.230998
Metropolis success percentage=54.090000
finished burn in
At beginning of Trace, LL=-5.477040
finished 10%
finished 20%
finished 30%
finished 40%
finished 50%
finished 60%
finished 70%
finished 80%
finished 90%
finished 100%
At end of trace, best LL=1.678685
Best Theta=
Metropolis success percentage=53.609600
writing, ntrace = 100001
writing, ntrace = 100001
```

For the sake of numerical efficiency 'Metropolis success percentage', should be in the range of 50%. If the efficiency is very near 100%, it suggests the step sizes might be too small to best explore the entire model-parameter space. If the efficiency is near zero, the step size might be too large and too few successful Metropolis steps will be taken. This affects only the efficiency, not the validity, so any success percentage between 10% and 90% should suffice. In the output, ''LL', refers to the log-likelihood. At the end of the burn-in, one hopes that best value of LL is not much lower than the best value found from the entire trace. If not, one should probably increase the value of MCMC\_NBURN. The values of ''Best Theta', refer to the point in the trace with the highest LL.

Information about the trace is found in the files located in the directory smooth\_data/mcmc\_trace/:

• Xtrace.txt: A list of the model-parameter values from the posterior sampling.

- trace.txt: A list of the scaled model-parameter values  $(\vec{\theta})$  from the posterior sampling.
- xbar\_thetabar.txt: The average of the parameter values  $(\vec{X})$ , and the scaled values  $(\vec{\theta})$  from the posterior.
- CovThetaTheta.txt: This gives the  $N_{par} \times N_{par}$  covariance matrix  $\langle \delta \theta_i \delta \theta_j \rangle$ , describing the size and shape of the points in the trace.
- CovThetaTheta\_eigenvalues.txt: That eigenvalues of that matrix
- CovThetaTheta\_eigenvecs.txt: The eigenvectors
- ResolvingPower.txt: An  $N_{pars} \times N_{obs}$  matrix describing the influence of each observable in resolving each model parameter.

# 7.4 Reviewing the MCMC Source Code

Finally, we review the source code in \${MY\_LOCAL}/software/main\_programs/mcmc\_main.cc:

```
int main(){
  CparameterMap *parmap=new CparameterMap();
  CSmoothMaster master(parmap);
  CMCMC mcmc(&master);
  //master.ReadCoefficientsAllY();
 master.TuneAllY();
  unsigned int Nburn=parmap->getI("MCMC_NBURN",1000); // Steps for burn in
  unsigned int Ntrace=parmap->getI("MCMC_NTRACE",1000); // Record this many points
  unsigned int Nskip=parmap->getI("MCMC_NSKIP",5); // Only record every Nskip^th point
  mcmc.PerformTrace(1,Nburn);
  CLog::Info("finished burn in\n");
  mcmc.PruneTrace(); // Throws away all but last point
  mcmc.PerformTrace(Ntrace, Nskip);
  mcmc.EvaluateTrace();
 mcmc.WriteTrace();
  return 0;
}
```

This is mostly self-explanatory. If one wishes to avoid writing out the trace, the line mcmc.WriteTrace can be deleted. If the User wishes to run the code in batch mode, the output can be directed to a file, mcmc\_log.txt, by adding the line

```
LogFileName mcmc_log.txt
```

to the parameter file parameters/mcmc\_parameters.txt. The line mcmc.EvaluateTrace() will evaluate the trace and calculate the resolving power and covariances.

## 8 Tutorial

#### 8.1 Overview

Template project directories and files are provided with the intention that the User will copy them to their own space, then use this as a foundation from which to embark on their own analysis. This directory includes information files, describing the parameter priors and the observables, that correspond to an artificial model that is also provided as a template. Working through the steps in this section constitutes a tutorial, both for running *Simplex Sampler* and for running *Smooth Emulator*.

This section describes the steps of how the User would

- 1. Copy the required files from the template directory to the User's space, and compile the main programs.
- 2. Set up the information files describing the priors and observable names.
- 3. Run Simplex Sampler to generate the model-parameter values at which the full model will be trained.
- 4. Run a full model to generate the observables for each of the full-model runs.
- 5. Tune Smooth Emulator and write the coefficients to file.
- 6. Run a program that prompts the User for the coordinates of a point in parameter space, then returns the emulator prediction with its uncertainty.

# 8.2 Installation and Compilation

Installation and compilation is described in Sec. 2. As was defined in that section, the tutorial will refer to three locations with the short hand:

\${GITHOME_BAND_SMOOTH}	Location of Git Repository, e.g. /Users/CarlosSmith/bandframework/software/SmoothEmulator
\${MY_LOCAL}	Can be placed anywhere. Executables are store in \$\{MY_LOCAL\}/bin and main programs, and source codes for main programs are found within \$\{MY_LOCAL\}/software/main_programs
\${MY_PROJECT}	Can be placed anywhere. Work spaces where parameter files, data, results and figures are created and stored. User may have several different such directories

At this point, the user should have established a personalized project directory by copying the \$\{\text{GITHOME\_BAND\_SMOOTH}\}/\text{templates/} \text{directory to a convenient location. For example,}

% cp -r \${GITHOME\_BAND\_SMOTH}/templates /Users/CarlosSmith/mysmoothy

In this case \${MY\_LOCAL} will now refer to Users/CarlosSmith/mysmoothy/mylocal and \${MY\_PROJECT} will now refer to a specific project at Users/CarlosSmith/mysmoothy/myproject.

The User should have also compiled the main libraries

```
${GITHOME_BAND_SMOOTH}/software% cmake . ${GITHOME_BAND_SMOOTH}/software% make
```

and the main programs,

```
${MY_LOCAL}/software% cmake .
${MY_LOCAL}/software% make
```

This will compile all the main source programs in \${MY\_LOCAL}/software/main\_programs/\*.cc and a "fake" full model to be used in the tutorial \${MY\_LOCAL}/software/fakefulmodels/fakerhic.cc. The repository was organized to encourage Users to edit any files in \${MY\_LOCAL}/. If the User wishes to restore any original files, a copy can be found at \${GITHOME\_BAND\_SMOOTH}/templates/mylocal.

Several executables should now appear in \${MY\_LOCAL}/bin/: simplex, smoothy\_tune, mcmc, smoothy\_testattrainingpts, smoothy\_testvsfullmodel, smoothy\_testvsfullmodelalt and fakerhic. The User might find it convenient to add \${MY\_LOCAL}/bin to their path. The reason these are compiled in the User's space, separate from the main libraries, is that the User may well wish to create their own main programs, and this arrangement allows the User to compile their own versions, while leaving the original programs from the templates directory and the lower-level source code unchanged.

## 8.3 Creating Necessary Info Files

The User will run the software from the \${MY\_PROJECT}/ directory. Before a User can run Simplex Sampler they must create information files that describe the model-parameter priors and list the observable names. Both files are need to be in the \${MY\_PROJECT} directory. The first file is smooth\_data/Info/modelpar\_info.txt, which describes the model parameters and their priors. For the purposes of this tutorial, a file already exists,

uniform	150	300
uniform	0.05	0.32
uniform	0.3	1.2
uniform	0.0	1.0
uniform	0.5	2.0
uniform	15.0	30.0
	uniform uniform uniform uniform	uniform 0.05 uniform 0.3

Thus, the model has six parameters. The second entry in each line is either uniform or gaussian. If the entry is uniform, the last two numbers represent the range of the uniform prior,  $x_{\min}$  and  $x_{\max}$ . If the second entry is gaussian the third entry represents the center of the Gaussian distribution and the fourth represents the width. For a full model, the User would replace this model with one appropriate for their own model.

The second file is smooth\_data/Info/observable\_info.txt. This describes output values from the model. In the template, the provided file is

meanpt_pion	100
meanpt_kaon	200
meanpt_proton	300
Rinv	1.0
v2	0.2
RAA	0.5

The first entry in each line simply provides the names of the observable which will be processed in the Bayesian analysis. The second entry is used by Smooth Emulator during tuning, but only if a Monte Carlo method is used, and then is only used to seed the Monte Carlo search. If the analytical method is used for tuning (which is recommended) this parameter is irrelevant (but still shoud be listed).

## 8.4 Running Simplex Sampler

Both Simplex Sampler and Smooth Emulator have options. These are provided in parameter files. For this tutorial, the provided parameter file is smooth\_data/smooth\_parameters/simplex\_parameters.txt. The provided file is

```
#LogFileName simplexlog.txt # comment out to direct output to screen
Simplex_TrainType 2 # Must be 1 or 2
Simplex_ModelRunDirName modelruns # Directory with training pt. info
```

Because the first line is commented, the output of  $Simplex\ Sampler\$ will be to the screen. Otherwise it would go to the specified file. By setting  $Simplex\_TrainType=1$ , the sampler will choose n+1 training points, where n=6 is the number of model parameters. Each point corresponds to the vertices of an n+1 dimensional simplex. Finally, the parameter  $Simplex\_ModelRunDirName$  is set to "modelruns". This informs  $Simplex\ Sampler\$ to write the coordinates of each training point and the corresponding observables in the directory  $smooth\_data/modelruns/$ . Here,  $Simplex\_TrainType=2$ , which adds points half-way between each pair of simplex points. These additional points are then moved outward and the original simplex points are brought inward. This method has precisely the number of training points as the number of coefficients necessary for a quadratic fit.

Now the user can run Simplex Sampler, which must be run from the project directory. The only output is the number of training points.

```
${MY_PROJECT}/rhic% ${MY_LOCAL}/bin/simplex
NTrainingPts=28
```

If one had set Simplex\_TrainType=1, only seven training points would have been created. The programs writes information about the training points in the smooth\_data/modelruns/ directory.

Changing into that directory, there should now be 28 sub-directories, corresponding to the 28 training points: modelruns/run0, modelruns/run1, modelruns/run2, modelruns/···. Each directory has one text file describing the training points. For example, the modelruns/run0/mod\_parameters.txt file might be

```
compressibility 190.282
etaovers 0.14892
initial_flow 0.664958
initial_screening 0.426807
quenching_length 1.16036
initial_epsilon 21.7424
```

This describes the six model parameters, which will serve as the input for the first full model run. The next step will be to run the full model for the parameters in each directory. Thus for Simplex\_Traintype=1, one would need 7 full-model runs, and for Simplex\_Traintype=2, one would need to do 28 full-model runs. The corresponding observables will be written in the files smooth\_data/modelruns/runI/obs.txt

## 8.5 Running the Fake Full Model

Once the training points have been generated, the user will run a full model based on the given structure, tailored to address their specific problem. For the tutorial, a fake full model is provided. It reads the model-parameter values in each smooth\_data/modelruns/runI/mod\_parameters.txt file and writes the corresponding observables in smooth\_data/modelruns/runI/obs.txt. The output should be as follows:

```
${MY_PROJECT}/rhic% ${MY_LOCAL}/bin/fakerhic
NTraining Pts=28
NPars=6
```

The output simply verifies the number of model parameters and the number of training points created by simplex.

Inspecting the smooth\_data/modelruns/run0/obs.txt file,

```
meanpt_pion
              418.821195
                           1.000000
meanpt_kaon
              715.592889
                           2.000000
meanpt_proton 1079.482871 3.000000
              5.004248
Rinv
                           0.010000
v2
              0.178353
                           0.002000
RAA
              0.553416
                           0.005000
```

The second entry of each line is the value of the specified observable for that specific training point. The last entry is the random uncertainty associated with the full model. This is only relevant if the model has random fluctuations, meaning the re-running the model at the same point might result

in different output. For this tutorial, the emulator will not consider such fluctuations (there is an emulator parameter that can be set to either consider the randomness or ignore it), so the third entry on each line is usually superfluous.

Additionally, fakerhic created a directory \${MY\_PROJECT}/fullmodel\_testdata/ which stores information about full-model runs at 50 randomly chosen points in the model-parameter space. These points are not used for tuning. This data can be used later to test the emulator.

## 8.6 Running Smooth Emulator

Before building and tuning the emulator, the User needs to edit one additional file, the parameter file that sets numerous options for *Smooth Emulator*. For the template used in this tutorial, that file is

```
#LogFileName smoothlog.txt # comment out for interactive running SmoothEmulator_LAMBDA 2.5 # smoothness parameter SmoothEmulator_MAXRANK 4 SmoothEmulator_ConstrainAO false SmoothEmulator_TrainingPts 0-27 SmoothEmulator_TestPts 1 SmoothEmulator_UsePCA false SmoothEmulator_TrainingFormat training_format_smooth #SmoothEmulator_TrainingFormat training_format_surmise SmoothEmulator_TrainingThetasFilename TrainingThetas.txt SmoothEmulator_TrainingObsFilename TrainingObs.txt
```

The parameters are described in detail in Sec. 5. The most relevant parameter is setting the smoothness parameter. Also, it is important to make sure that SmoothEmulator\_TrainingPts is set to the correct number of training points.

Now, running smoothy\_tune, produces the following output,

```
${MY_PROJECT}/rhic% ${MY_LOCAL}/bin/smoothy_tune
---- Tuning for meanpt_pion ----
---- Tuning for meanpt_kaon ----
---- Tuning for meanpt_proton ----
---- Tuning for Rinv ----
---- Tuning for v2 ----
---- Tuning for RAA ----
.
```

The program generates Taylor coefficients which are saved in the coefficients/directory. Each observable has its own sub-directory with its name. In this case, smoothy\_tune created the directories, coefficients/rhic/RAA, coefficients/Rinv, coefficients/menapt\_kaon, coefficients/meanpt\_pion, coefficients/meanpt\_proton and coefficients/v2. Within each of these sub-directories smoothy\_tune

created files meta.txt, ABest.txt and BetaBest.txt. The number or parameters, the maximum rank of the Taylor expansion and the overall number of Taylor coefficients are give in meta.txt. The file ABest.txt provides the actual coefficients of the Taylor expansion, and BetaBest.txt gives an array used to calculate the uncertainty.

To get an idea of how one might build one's own main program to access the capabilities used above, the source code for  $MY_LOCAL$ /software/main\_programs/smoothy\_tune\_main.cc is:

```
#include "msu_smoothutils/parametermap.h"
#include "msu_smooth/master.h"
#include "msu_smoothutils/log.h"
using namespace std;
int main(){
   NMSUUtilsCparameterMap *parmap=new CparameterMap();
   NBandSmooth::CSmoothMaster master(parmap);
   master.TuneAllY();
   master.WriteCoefficientsAllY();
   master.TestAtTrainingPts();
   return 0;
}
```

Hopefully, the User will find this and the other main-program source codes to be fairly self-explanatory. Nonetheless, detailed explanations can be found in Sec. 5. Given that the tuning is very fast, there is little need to write the coefficients as any subsequent use of the emulators can simply repeat the tuning, rather than reading in the coefficients.

# 9 Testing the Emulator at the Training Points

The last line in the source code above tells  $Smooth\ Emulator$  to evaluate the emulator at the training points. The remainder of the outpus is:

```
--- Y_train
               Y_emulator
                              Sigma_emulator ----
----- itrain=0 ------
Y[0] = 3.999e + 02 = ? 3.999e + 02
                               +/-
                                    1.11725e-04
Y[1]= 6.758e+02 =? 6.758e+02
                                    1.71543e-04
                               +/-
Y[2] = 1.090e+03 =? 1.090e+03
                               +/-
                                    3.46421e-04
Y[3] = 4.916e+00 =? 4.916e+00
                               +/-
                                    1.93211e-06
                   3.691e-01
Y[4] = 3.691e-01 = ?
                               +/-
                                    2.94503e-07
Y[5] = 1.980e-01 =?
                   1.980e-01
                               +/-
                                    5.60746e-07
----- itrain=1 -----
Y[0] = 4.686e+02 = ? 4.686e+02
                              +/- 7.64376e-05
Y[1]= 7.596e+02 =? 7.596e+02
                               +/-
                                    1.17363e-04
Y[2] = 1.107e+03 =? 1.107e+03 +/-
                                    2.37007e-04
```

```
Y[3]= 5.013e+00 =?
                                   1.32187e-06
                   5.013e+00
                               +/-
Y[4] = 4.705e-01 = ?
                   4.705e-01
                               +/-
                                   2.01488e-07
Y[5] = 3.210e-01 = ?
                   3.210e-01
                                   3.83641e-07
----- itrain=2 -----
Y[0] = 4.721e+02 = ? 4.721e+02
                               +/-
                                   1.14852e-04
Y[1]= 7.837e+02 =?
                   7.837e+02
                                   1.76346e-04
                              +/-
Y[2]= 1.129e+03 =?
                   1.129e+03
                              +/-
                                   3.56118e-04
----- itrain=27 -----
Y[0] = 4.527e+02 = ? 4.527e+02
                              +/-
                                   3.37022e-05
Y[1]= 7.138e+02 =? 7.138e+02
                              +/-
                                   5.17468e-05
Y[2] = 1.075e+03 =? 1.075e+03
                              +/-
                                   1.04499e-04
Y[3] = 6.153e+00 =? 6.153e+00
                              +/-
                                   5.82829e-07
Y[4] = 3.235e-01 = ? 3.235e-01
                               +/-
                                   8.88382e-08
Y[5]=-2.208e-02 =? -2.208e-02
                              +/-
                                   1.69151e-07
```

The observables,  $Y[0] \cdots Y[5]$  should be identical and the uncertainties at the training points should be zero. The fact that the uncertainties are not exactly zero derives from the numerical accuracy of the linear algebra routines. For this model, the random uncertainties were set to zero. If non-zero random uncertainties had been applied (would require adjusting the last column in each of the  $smooth_data/modelruns/run*/obs.txt$  files) then the emulator would not have exactly reproduced the training values.

# 10 Generating Emulated Observables at Given Points

Finally, now that the emulator is tuned, one may wish to generate emulated values for the observables for specified points in model-parameter space. A sample program, \${MY\_LOCAL}/bin/smoothy\_calcobs is provided to illustrate how this can be accomplished. If one invokes the executable, using the same parameters as those used by smoothy\_tune, the User is prompted to enter the coordinates of a point in model-parameter space, after which smoothy\_calcobs prints out the observables. In this case, for the case where compressibility=225, etaovers=0.2, initial\_flow=0.5, initial\_screening=0.6, quenching\_length=1.2 and initial\_epsilon=25.0

\${MY\_PROJECT}/rhic% \${MY\_LOCAL}/bin/smoothy\_calcobs
Prior Info

#	ParameterName	Type	Xmin_or_Xbar	Xmax_or_SigmaX
0:	compressibility	uniform	150	300
1:	etaovers	uniform	0.05	0.32
2:	<pre>initial_flow</pre>	uniform	0.3	1.2
3:	initial_screening	uniform	0	1
4:	quenching_length	uniform	0.5	2
5:	initial_epsilon	uniform	ı 15	30
п.	7 6			

Enter value for compressibility:

```
225
Enter value for etaovers:
Enter value for initial_flow:
Enter value for initial_screening:
Enter value for quenching_length:
Enter value for initial_epsilon:
---- EMULATED OBSERVABLES -----
meanpt_pion = 451.556 + / - 9.31878
meanpt_kaon = 751.461 + - 14.3082
meanpt_proton = 903.253 +/- 28.8944
Rinv = 5.13761 + /- 0.161154
v2 = 0.247525 + - 0.0245641
RAA = 0.0800345 +/- 0.046771
Of course, it is unlikely the User will wish to enter model parameters interactively as was done
above. To incorporate Smooth Emulator into other programs, the User should inspect the main
programs, e.g. $\{MY_LOCAL\}/software/main_programs/smoothy_calcobs_main.cc. The User can
then design their own program based on this source code, and compile and link it by editing
${MY_LOCAL}/software/main_programs/CMakeLists.txt. By editing the CMake file, replacing
the lines unique to smoothy_calcobs, one can easily compile new executables based on the User's
main programs. To understand what might be involved, the source code in
${MY_LOCAL}/software/main_programs/smoothy_calcobs_main.cc is
#include "msu_smoothutils/parametermap.h"
#include "msu_smooth/master.h"
#include "msu_smoothutils/log.h"
using namespace std;
int main(){
NBandSmooth::CSmoothMaster master;
master.ReadCoefficients();
 //master.TuneAllY();
 //Create model parameter object to store information about a single set of model parameter
 NBandSmooth::CModelParameters *modpars=new NBandSmooth::CModelParameters(); // contains in
 modpars->priorinfo=master.priorinfo;
 // Print out the prior information
 master.priorinfo->PrintInfo();
 // Prompt user for model parameter values and enter them into the modpars object
 vector<double> X(modpars->NModelPars);
 for(unsigned int ipar=0;ipar<modpars->NModelPars;ipar++){
```

```
cout << "Enter value for " << master.priorinfo->GetName(ipar) << ":\n";
    cin >> X[ipar];
}
modpars->SetX(X);

// Calc Observables
NBandSmooth::CObservableInfo *obsinfo=master.observableinfo;
vector<double> Y(obsinfo->NObservables);
vector<double> SigmaY(obsinfo->NObservables);
master.CalcAllY(modpars,Y,SigmaY);
cout << "---- EMULATED OBSERVABLES -----\n";
for(unsigned int iY=0;iY<obsinfo->NObservables;iY++){
    cout << obsinfo->GetName(iY) << " = " << Y[iY] << " +/- " << SigmaY[iY] << endl;
}
return 0;
}</pre>
```

The above illustrated how one can write a code that

- a) Reads the parameter file.
- b) Creates a *master* emulator file.
- c) Read the Taylor coefficients that were written when the emulator was tuned. (One could have commented out this line and retuned instead)
- d) Creates a model-parameters object, modpars, that stores the coordinates of the model-parameter point
- e) Reads in the model parameters interactively
- f) Calculates the observables from the emulator
- g Prints out the emulated observable and the uncertainty for for the emulator

# 10.1 Exploring the Posterior via Markov-Chain Monte-Carlo

Given the experimental information, which is stored in project directory in smooth\_data/Info/experimental\_info one can then use the tuned emulator to explore the posterior likelihood through MCMC, which works via a Metropolis algorithm. The file smooth\_data/Info/experimental\_info.txt provided in the template is:

meanpt_pion	481.179	20	0.0
meanpt_kaon	757.872	30	0.0
meanpt_proton	1113.3	40	0.0
Rinv	6.27842	0.3	0.0
v2	0.382973	0.1	0.0
RAA	0.558367	0.1	0.0

The first column is the list of observable names, which should be identical to those listed in  $smooth_data/Info/observable_info.txt$ . The second and third columns lists the experimental measurement,  $Y_a$ , and the experimental uncertainty,  $\sigma_a^{exp}$ . The last column lists the additional uncertainty due to errors,  $\sigma_a^{theory}$ , i.e. missing physics, in the theoretical model. For the purposes of comparing theory to data, only the combination  $(\sigma_a^{exp})^2 + (\sigma_a^{exp})^2$  comes into play, because this combination appears in the likelihood for the posterior,

$$\mathcal{L}(\vec{\theta}) = \prod_{a} \frac{1}{\sqrt{2\pi(\sigma_a^{\text{tot}})^2}} \exp\left\{-\frac{(Y_a(\vec{\theta}) - Y_a^{\text{exp}})^2}{2(\sigma_a^{\text{tot}})^2}\right\}$$

$$(\sigma_a^{\text{tot}})^2 = (\sigma_a^{\text{exp}})^2 + (\sigma_a^{\text{theory}})^2 + (\sigma_a^{\text{emu}})^2.$$
(10.1)

Whereas the emulator uncertainty,  $\sigma_a^{\text{emu}}$ , depends on the location in parameter space,  $\vec{\theta}$ , the other two contributions are assumed to be independent of  $\vec{\theta}$ .

There are special parameters for the MCMC. These are stored in smooth\_data/Info/mcmc\_parameters.txt. For the tutorial template, that file is

# This is for the MCMC search of parameter space
# (not for the emulator tuning)
MCMC\_LANGEVIN false
MCMC\_METROPOLIS\_STEPSIZE 0.05
MCMC\_LANGEVIN\_STEPSIZE 0.5
MCMC\_NBURN 100000
MCMC\_NTRACE 100000
MCMC\_NSKIP 5
RANDY\_SEED 12345

The first parameter, MCMC\_LANGEVIN should be set to false, as the Langevin MCMC (as opposed to the Metropolis version) is under development. The Metropolis stepsize should be adjusted so that the Metropolis success rate is approximately one half. The success rate prints out when the mcmc code runs. If the success rate is anywhere between 20 and 80%, this should be fine. But, if the rate is close to zero or 100%, the efficiency of the procedure suffers. It is recommended to run the MCMC code with a modest number of steps, then adjust the stepsize accordingly.

The parameter MCMC\_NBURN sets the number of Metropolis steps to be used in the "burn-in" stage, i.e. before one begins to store the trace. The number of elements to store in the trace in MCMC\_NTRACE, and MCMC\_NSKIP sets the number of steps to skip before storing a new point in the trace. Thus, if MCMC\_NTRACE is one million and if MCMC\_NSKIP=5, then the procedure will perform 5 million steps, and store every fifth one, leading to one million stored points in the trace. Finally, RANDY\_SEED sets the random number seed.

Running the MCMC program gives the following output:

\${MY\_PROJECT}/rhic% \${MY\_LOCAL}/bin/mcmc
At beginning of Trace, LL=-68.728122
At end of trace, best LL=1.223190
Best Theta=

```
Metropolis success percentage=55.010000
finished burn in
At beginning of Trace, LL=-3.055348
finished 10%
finished 20%
finished 30%
finished 40%
finished 50%
finished 60%
finished 70%
finished 80%
finished 90%
finished 100%
At end of trace, best LL=1.375865
Best Theta=
0.236218  0.138233  0.183291  0.217895
                                   0.033619
                                            0.194723
Metropolis success percentage=57.355500
writing Theta values, ntrace = 100001
writing X values, ntrace = 100001
```

Here best LL refers to the log-likelihood and Best Theta refers to the value of  $\vec{\theta}$  that gave the maximum log-likelihood. Values of Best Theta and best LL are given after the burn-in and after the trace. One can read the source file,  ${MY\_LOCAL}/software/main\_programs/mcmc\_main.cc$ 

The trace is written to mcmc\_trace/trace.txt. It is in the format

```
theta_1 theta_2 theta_3 theta_4 ... theta_1 theta_2 theta_3 theta_4 ... theta_1 theta_2 theta_3 theta_4 ...
```

If MCMC\_NTRACE is set to a million, there would be a million lines in the file. The program also calculated various covariances:  $\langle\langle\delta\theta_i\delta\theta_j\rangle\rangle$ ,  $\langle\langle\delta\theta_i\delta Y_a\rangle\rangle$ , and  $\langle\langle\delta Y_a\delta Y_b\rangle\rangle$ . The quantities  $\langle\langle....\rangle\rangle$  refer to averages over the posterior, i.e. averages over the trace. The eigenvalues and eigenvectors of  $\langle\langle\delta\theta_i\delta\theta_j\rangle\rangle$  are also recorded. Hopefully, the User will find the file names in mcmc\_trace/ to be self-explanatory.

As described in Sec. 9 the covariances in the posterior can also be used to quantify the resolving power of specific observables to constrain specific parameters. One such measure is

$$\mathcal{R}_{ia} \equiv \frac{d\langle\langle\theta_i\rangle\rangle}{dY_a^{\text{exp}}} \langle\delta Y_a^2\rangle^{1/2}.$$
 (10.2)

This quantifies how the posterior value of  $\theta_i$  changes as the experimental value changes if the experimental value,  $Y_a^{\text{exp}}$ , changes an amount characteristic of the variance of  $Y_a$  across the prior. Given that there may not be a set of training points calculated uniformly across the prior, the final factor is estimated as described in Sec. 9. Higher values of  $\mathcal{R}_{ia}$  for different a demonstrate the relative contributions of different observables a to constrain a model parameter i. The resolving power matrix is written to mcmc\_trace/ResolvingPower.txt.

## 10.2 Making Plots

Three python scripts (using MATPLOTLIB) are provided to provide graphical insight into the posterior likelihood, into the resolving power and for viewing how the emulator uncertainty compares to the discrepancies between full-model runs (not used for tuning) and emulated value. One can visit the \${MY\_PROJECT}/figs/ directory and peruse the file directions.txt for more detailed instructions of how to create the plots below. First, one must create two data files. These provide the names of observables and model parameters to be used by the plots.

The two files are given the same names as two files in <code>smooth\_data/Info/</code>. The first is <code>\${MY\_PROJECT}/figs/mooth\_data/Info/</code> directory in that it has only three columns, though the first column is identical. The provided file is

```
$\kappa$
compressibility
                           uniform
                                      $\eta/s$
                           uniform
etaovers
                                      Init.Flow
initial_flow
                           uniform
initial_screening
                           uniform
                                      Screening
quenching_length
                           uniform
                                      $\lambda_{\rm quench}$
initial_epsilon
                                      $\epsilon_0$
                           uniform
\end{verbatime}}
```

As one can see, the last column is used by MATPLOTLIB to label axes. The middle column is n

The second required file is  ${\t \ \}MY\_PROJECT\}/figs/observable\_info.txt\}$ , and the prov  ${\t \}$ 

\begin{verbatim}

Rinv  $R_{\text{inv}}$ 

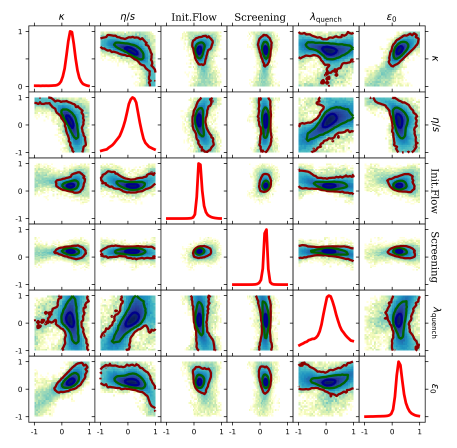
v2 \$v\_2\$ RAA \$R\_{AA}\$

Again, the first column is identical to that in smooth\_data/Info/ and the second is used for labeling.

To graph the posterior likelihood, first be sure to run the mcmc program, then change into the figs/posterior directory and enter the command:

```
${MY_PROJECT}/figs/posterior% ln -s ../../smooth_data/mcmc_trace/trace.txt . ${MY_PROJECT}/figs/posterior% python3 posterior.py
```

One can replace "ln -s" with "cp" or "mv". The script should create a file posterior.pdf, which looks like:



Projections for the posterior likelihood from the MCMC trace. The contour lines represent  $1-\sigma$ ,  $2-\sigma$  and  $3-\sigma$  likelihoods.

The likelihood is projected for individual model parameters, or for pairs. The plot is in terms of the scaled variables,  $\theta_i$ . To translate to the true model-parameter ranges, one can look at the smooth\_data/Info/modelpars\_info.txt file, which gives the prior ranges of the model parameters before they are scaled to the -1 to 1 range. The file figs/directions.txt shows how the User can alter plot. For example there is a line in the python script, ParsToPlot=[1,2,3,4,5,0], which the User can edit to change the ordering of the model-parameters, and to choose which model parameters are considered.

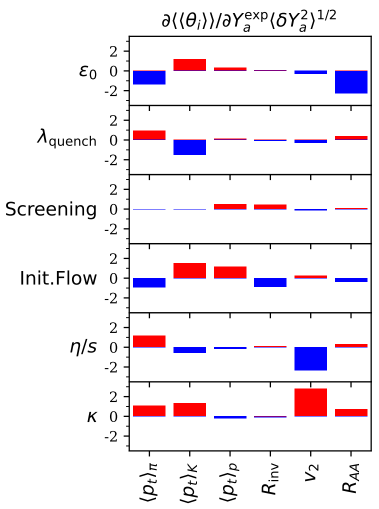
Similarly, one can plot the resolving power. The User can visit the directory figs/resolvingpower/. The User should copy the files mcmc\_trace/ResolvingPower.txt and smooth\_data/Info/modelpar\_info.txt to this directory, editing the smooth\_data/Info/modelpar\_info.txt as was done for the posterior visualization figures described above. The User must also copy over the smooth\_data/Info/observable\_info.file and edit it in a similar fashion. In the template file is

meanpt_pion	<pre>\$\langle p_t\rangle_{\pi}\$</pre>
meanpt_kaon	<pre>\$\langle p_t\rangle_{K}\$</pre>
meanpt_proton	<pre>\$\langle p_t\rangle_{p}\$</pre>
Rinv	\$R_{\rm inv}\$
v2	\$v_2\$
R.A.A	\$R. {AA}\$

The first column should be exactly the same as the first line in the original file. After running the mcmc program, the figure can be produced via the command

 $\$  WY\_PROJECT}/figs/resolving power% ln -s ../../smooth\_data/mcmc\_trace/Resolving Power.txt .  $\$  WY\_PROJECT}/figs/resolving power% python 3 RP.py

The figure should look like:



Resolving Power. Red bars represent positive correlations with  $Y_a^{\text{exp}}$  and  $\theta_i$ . Larger bars suggest that the particular observable contributes more to the constraint of the particular model parameter.

The third provided python script is in figs/YvY/ and it compares full-model runs (not used for tuning) to the emulator. This is useful for seeing whether the emulator's error estimates are reasonable. First, one must run the program that writes out the emulator predictions for the full-model runs. This is accomplished by the command:

\${MY\_PROJECT}% smoothy\_testvsfullmodelalt

meanpt\_pion: 27 out of 50 points within 1 sigma
meanpt\_kaon: 41 out of 50 points within 1 sigma
meanpt\_proton: 26 out of 50 points within 1 sigma

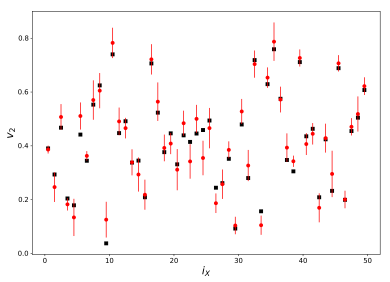
Rinv: 32 out of 50 points within 1 sigma v2: 39 out of 50 points within 1 sigma RAA: 29 out of 50 points within 1 sigma

If the uncertainty were perfectly stated, 68% of the points would be within one standard deviation.

In this case the fraction was higher, which suggests that the uncertainty was somewhat overstated. To make the plot, first change into the figs/YvsY/ directory, and enter

```
${MY_PROJECT}/figs/YvsY% ln -s ../../fullmodel_testdata .
${MY_PROJECT}/figs/YvsY% python3 YvsY.py
Observable names:
['meanpt_pion', 'meanpt_kaon', 'meanpt_proton', 'Rinv', 'v2', 'RAA']
Enter iY: 4
39 of 49 points within 1 sigma
```

The script will prompt the User for which observable to consider. In this case, choose 0-5 for the six possible observables. In this case '4' was entered and the chosen observable was the elliptic flow v2.



Comparison of full-model values (black squares) for 50 points in parameter space to emulator values (red circles). The uncertainties are solely those associated with the emulation. If the uncertainties were accurately expressed, 68% of the points would lie within the uncertainty intervals.

For this example the full-model test data was stored in a few files in the smooth\_data/fullmodel\_testdata/ directory, consistent with the SURMISE format. Another option would be to use full-model run data stored in the same format and location as the training data. In that case one would invoke the program smoothy\_testvsfullmodel (with the alt) and the testing runs would be denoted by the SmoothEmulator\_TestPts parameter set in the smooth\_data/smooth\_parameters/emulator\_parameters.txt file. Those model runs should be chosen separately from those used for training, i.e. those denoted by the SmoothEmulator\_TrainingPts parameter.

## 10.3 PCA Analysis

The PCA functionality needs further testing and is not included in the tutorial at this time.

# 9 Underlying Theory of Smooth Emulator

The choice of model emulators,  $E(\vec{\theta})$ , depends on the prior understanding of the model being emulated,  $M(\vec{\theta})$ . If one knows that a function is linear, then a linear fit is clearly the best choice. Whereas to reproduce lumpy features, where the lumps have a characteristic length scale, Gaussian process emulators are highly effective. The quality of an emulator can be assessed through the following criteria:

- $E(\vec{\theta}_t) = M(\vec{\theta}_t)$  at the training points,  $\vec{\theta}_t$ .
- The emulator should reasonably reproduce the model away from the training points. This should hold true for either interpolation or extrapolation.
- The emulator should reasonably represent its uncertainty
- A minimal number of training points should be needed
- The method should easily adjust to larger numbers of parameters,  $\theta_i$ ,  $i=1\cdots N$
- The emulator should not be affected by unitary transformations of the parameter space
- The emulator should be able to account for noisy models
- Training and running the emulator should not be numerically intensive

Here the goal is to focus on a particular class of functions: functions that are *smooth*. Smoothness is a prior knowledge of the function. It is an expectation that the linear terms of the function are likely to provide more variance than the quadratic contributions, which are in turn likely to be more important than the cubic corrections, and so on.

#### 9.1 Mathematical Form of Smooth Emulator

To that end the following form for  $E(\vec{\theta})$  is chosen,

$$E(\vec{\theta}) = \sum_{\vec{n}, \text{s.t. } K(\vec{n}) \leq K_{\text{max}}} d_{\vec{n}} f_{K(\vec{n})}(|\vec{\theta}|) A_{\vec{n}} \left(\frac{\theta_1}{\Lambda}\right)^{n_1} \left(\frac{\theta_2}{\Lambda}\right)^{n_2} \cdots \left(\frac{\theta_N}{\Lambda}\right)^{n_N}.$$
(9.1)

Each term has a rank  $K(\vec{n}) = n_1 + n_2 + \cdots + n_N$ . If f is constant, the rank of that term corresponds to the power of  $|\vec{\theta}|/\Lambda$ . All terms are included up to a given rank,  $K_{\text{max}}$ . The coefficients A are stochastically distributed. The coefficients  $d_{\vec{n}}$  will ensure that the variance is independent of the direction of  $\vec{\theta}$ , with the constraint that  $d_{K,0,0,\dots} = 1$ . The function  $f_K(|\vec{\theta}|)$  provides the freedom to alter how the behavior depends on the distance from the origin,  $|\vec{\theta}|$ , and on the rank, K. Given that the variance of  $A_{\vec{n}}$  can be changed,  $f_{K=0}(|\vec{\theta}| = 0)$  is also set to unity for all K without loss of generality. For each combination  $\vec{n}$ , the prior probability for any the A coefficients is given by

$$p(A_{\vec{n}}) = \frac{1}{\sqrt{2\pi\sigma_{K(\vec{n})}^2}} e^{-A_{\vec{n}}^2/2\sigma_{K(\vec{n})}^2},$$

$$\langle A_{\vec{n}}^2 \rangle = \sigma_{K(\vec{n})}^2.$$
(9.2)

The variance,  $\sigma_K^2$ , is allowed to vary as a function of K.

The parameter  $\Lambda$  will be referred to as the *smoothness parameter*. Here, we assume that all parameters have a similar range, of order unity, e.g.  $-1 < \theta_i < 1$ . Thus, the relative importance of each term Eq. (9.1) falls with increasing rank, K, as  $(1/\Lambda)^K$ . For now, the smoothness parameter is fixed by prior knowledge, i.e. one chooses higher values of  $\Lambda$  if one believes the function to be close to linear.

First, we consider the variance of the emulator at a given point,  $\vec{\theta}$ . Requiring that the variance is independent of the direction of  $\vec{\theta}$  will fix  $d_{\vec{n}}$ . For example, transforming  $\theta_1$  and  $\theta_2$  to parameters  $(\theta_1 \pm \theta_2)/\sqrt{2}$  should not affect the accuracy or uncertainty of the emulator.

At  $|\vec{\theta}| = 0$  the only term in Eq. (9.1) that contributes to the variance is the one K = 0 term. Averaging over the A coefficients, which can be either positive or negative with equal probability,

$$\langle E(\vec{\theta}) \rangle = 0, \tag{9.3}$$

where the averaging refers to an average over the A coefficients. At the origin,  $|\vec{\theta}| = 0$ , the variance of E is

$$\langle E(\theta_1 = \theta_2 = \cdots \theta_N = 0)^2 \rangle = d_{n_i=0}^2 \sigma_{K=0}^2 f_{K=0}^2 (\vec{\theta} = 0).$$
 (9.4)

Choosing  $f_{K=0}(0) = 1$  and  $d_{n_i=0} = 1$ , the variance of E is indeed  $\sigma_0^2$ . The variance at some point  $\vec{\theta} \neq 0$  is

$$\langle E^2(\vec{\theta}) \rangle = \sum_{\vec{n}} f_K^2(|\vec{\theta}|) \sigma_{K(\vec{n})}^2 d_{\vec{n}}^2 \left( \frac{\theta_1^{2n_1}}{\Lambda^2} \right) \left( \frac{\theta_2^{2n_2}}{\Lambda^2} \right) \cdots \left( \frac{\theta_N^{2n_N}}{\Lambda^2} \right). \tag{9.5}$$

If  $\langle E^2 \rangle$  is to be independent of the direction of  $\vec{\theta}$ , the sum above must be a function of  $|\vec{\theta}|^2$  only. This requires the net contribution from each rank, K to be proportional to  $|\vec{\theta}|^{2K}$  multiplied by some function of K. Using the fact that

$$(\vec{\theta}_a \cdot \vec{\theta}_b)^K = \sum_{\vec{n}, s.t. \sum_i n_i = K} \frac{K!}{n_1! \cdots n_N!} (\theta_{a1}\theta_{b1})^{n_1} \cdots (\theta_{aN}\theta_{bN})^{n_N}, \tag{9.6}$$

one can see that if the sum is to depend only on the norm of  $\vec{\theta}$ ,

$$d_{\vec{n}}^2 = \frac{K(\vec{n})!}{n_1! n_2! \cdots n_N!}.$$
 (9.7)

The factor of K! in the numerator was chosen to satisfy the condition that  $d_{K,0,0,0}=1$ .

One can now calculate the correlation between the emulator at two different points, averaged over all possible values of A,

$$\langle E(\vec{\theta}_a)E(\vec{\theta}_b)\rangle = \sum_{K=0}^{K_{\text{max}}} \sigma_K^2 f_K^2(|\vec{\theta}|) \left(\frac{\vec{\theta}_a \cdot \vec{\theta}_b}{\Lambda^2}\right)^K.$$
(9.8)

Requiring  $f(|\theta| = 0) = 1$  gives

$$\langle E^2(\vec{\theta} = 0) \rangle = \sigma_0^2, \tag{9.9}$$

and for  $\vec{\theta}_a = \vec{\theta}_b$ ,

$$\langle E^2(\vec{\theta} = 0) \rangle = \sum_{K=0}^{K_{\text{max}}} \sigma_K^2 f_K^2(|\vec{\theta}|) \left(\frac{|\vec{\theta}|^2}{\Lambda^2}\right)^K.$$
 (9.10)

To this point, the form is completely general once one requires that the variance above is independent of the direction of  $\vec{\theta}$ . I.e. the function  $f_K(\vec{\theta})$  could be any function satisfying the constraint,  $f_K(0) = 1$ , and  $\sigma_K^2$  could have any function of K. Below, we illustrate how different choices for f or for  $\sigma_K$  affect the emulator by comparing several variations. First, we define the default form.

#### 9.2 Alternate Forms

As stated above, once the form is to provide variances that are independent of the direction of  $\vec{\theta}$ , the general form going forward is

$$E(\vec{\theta}) = \sum_{\vec{n}, \text{s.t. } K(\vec{n}) < K_{\text{max}}} f_K(|\vec{\theta}|) \left( \frac{K(\vec{n})!}{n_1! \cdots n_N!} \right)^{1/2} A_{\vec{n}} \left( \frac{\theta_1}{\Lambda} \right)^{n_1} \left( \frac{\theta_2}{\Lambda} \right)^{n_2} \cdots \left( \frac{\theta_N}{\Lambda} \right)^{n_N},$$

$$(9.11)$$

$$P(\vec{A}_n) = \frac{1}{(2\pi\sigma_K^2)^{1/2}} e^{-|A_{\vec{n}}|^2/2\sigma_K^2}.$$

Variations from the general form involve adjusting either the K-dependence of the  $|\vec{\theta}|$ -dependence of  $f_K(|\vec{\theta}|)$ , or adjusting the K-dependence of  $\sigma_K$ .

#### Default Form

Here, we assume  $f_K(|\vec{\theta}|)$  is independent of  $|\vec{\theta}|$ , and that  $\sigma_K$  is independent of K. Further, the K-dependence of  $f^2$  is assumed to be 1/K!. With this choice

$$E(\vec{\theta}) = \sum_{\vec{n}, K(\vec{n}) \le K_{\text{max}}} \frac{1}{\Lambda^K} \frac{A_{\vec{n}}}{\sqrt{n_1! n_2! \cdots n_N!}} \theta_1^{n_1} \cdots \theta_N^{n_N},$$

$$P(A_{\vec{n}}) \sim e^{-A_{\vec{n}}^2/2\sigma^2}.$$
(9.12)

With this form the variance increases with  $|\vec{\theta}|$ ,

$$\langle E^2(\vec{\theta}) \rangle = \sigma^2 e^{|\vec{\theta}|^2/\sigma^2}.$$
 (9.13)

If the function is trained in a region where the function is linear, the emulator's extrapolation outside the region will continue to be follow the linear behavior, albeit with variation from the higher order coefficients.

The choice of  $f_K^2 = 1/K!$  ensures that the sum defining  $E(\vec{\theta})$  converges as a function of K as long as  $K_{\text{max}}$  is rather large compared to  $|\vec{\theta}|/\Lambda$ .

#### Variant A: Letting $\sigma_K$ have a K dependence

One reasonable alteration to the default choice might be to allow the K=0 term to take any value, i.e.  $\sigma_{K=0}=\infty$ , while setting all the other  $\sigma_K$  terms equal to one another. This would make sense if our prior expectation of smoothness meant that we expect the K=2 terms to be less important than the K=1 terms, by some factor  $|\vec{\theta}|/\Lambda$ , but that the variation of the K=1 term is unrelated to the size of the K=0 term. This would make the emulator independent of redefinition of the emulated function by adding a constant. This may well be a reasonable choice for many circumstances.

## Variant B: Suppressing correlations for large $\Delta \vec{\theta}$

This form for f causes correlations to fall for points far removed from one another.

$$f_K(|ec{ heta}|) = rac{1}{\sqrt{K!}} \left\{ \sum_{K=0}^{K_{ ext{max}}} rac{1}{\sqrt{K!}} \left( rac{|ec{ heta}|^2}{2\Lambda^2} 
ight)^K 
ight\}^{-1/2}.$$

In the limit that  $K_{\text{max}} \to \infty$  the form is a simple exponential,

$$f_K(|\vec{\theta}|)\Big|_{K_{\text{max}}\to\infty} = \frac{1}{\sqrt{K!}} e^{-|\vec{\theta}|^2/2\Lambda^2}.$$
 (9.14)

With this form the same-point correlations remain constant over all  $\vec{\theta}$ ,

$$\langle E(\vec{\theta})E(\vec{\theta})\rangle = \sigma^2,$$
 (9.15)

while the correlation between separate positions fall with increasing separation. This is especially transparent for the  $K_{\text{max}} \to \infty$  limit,

$$\langle E(\vec{ heta}_a) E(\vec{ heta}_b) 
angle_{K_{
m max} 
ightarrow \infty} = \sigma^2 e^{-|\vec{ heta}_a - \vec{ heta}_b|^2/2\Lambda^2}.$$

In this limit one can also see that

$$\langle [E(\vec{\theta}) - E(\vec{\theta'})]^2 \rangle_{K_{\text{max}} \to \infty} = 2\sigma^2 \left( 1 - e^{|\vec{\theta} - \vec{\theta'}|^2 / 2\Lambda^2} \right). \tag{9.16}$$

If one trains such an emulator in one region, then extrapolates to a region separated by  $|\vec{\theta} - \vec{\theta'}| >> \Lambda$ , the average predictions will return to zero. Thus, if the behavior would appear linear in some region the emulator's distribution of predictions far away (extrapolations) would center at zero. This behavior is similar to a Gaussian-process emulator.

### Variant C: Eliminating the 1/K! weight

Clearly, eliminating the 1/K! weights in  $f_K$  would more emphasize the contributions from larger K. But, for  $|\vec{\theta}| > \Lambda$  the contribution to the variance would increase as K increases and the sum would not converge if  $K_{\text{max}}$  were allowed to approach infinity. An example of a function that expands without factorial suppression is  $1/(1-x) = 1 + x + x^2 + x^3 \cdots$ , which diverges as  $x \to 1$ . If such behavior is not expected, then this choice would be unreasonable.

# 9.3 Exact Solution for the Most Probable Coefficients and the Uncertainty

Here, we demonstrate how one can solve for the set of most probable coefficients,  $A_{\vec{n}}$ , given the training values,  $y_t(\vec{\theta_t})$ , where  $\vec{\theta_t}$  are the points at which the emulator was trained. Again, we

consider  $N_{\text{train}}$  to be the number of training points and  $N_{\text{coef}}$  to be the number of coefficients, which is much larger than  $N_{\text{train}}$ . Given the coefficients,  $\vec{A}_i$  for  $i > N_{\text{train}}$ , the first coefficients,  $i = 1 \cdots N_{\text{train}}$ , are found by solving the linear equations for  $A_a, a \leq N_{\text{train}}$ . For shorthand, we define the function  $T_a(\vec{\theta})$ ,

$$E(\vec{\theta}) = \sum_{i=1}^{N_{\text{coeff}}} A_i \mathcal{T}_i(\vec{\theta}), \qquad (9.17)$$

The functions  $\mathcal{T}_i(\vec{\theta})$  reference the factors in Eq. (9.12) if the default form is used. Evaluated at the  $N_{\text{train}}$  training points, we define a  $N_{\text{train}} \times N_{\text{coef}}$  matrix,

$$T_{ai} \equiv \mathcal{T}_i(\vec{\theta}_a),$$
 (9.18)

where  $\vec{\theta}_a$  labels the training points,  $0 < a \le N_{\text{train}}$ , and  $N_{\text{train}} << N_{\text{coef}}$ .

#### 9.3.1 Finding the Optimum Coefficients and Uncertainty

The goal is to find the coefficients,  $A_{i>N_{\text{train}}}$ , that maximizes the probability given the constraints of matching the training data. If the training data are to be exactly matched, the differential probability of a given set of coefficients

$$dP(\vec{A}, \sigma_A, \Lambda) \sim \frac{1}{(2\pi\sigma_A^2)^{N_{\text{coef}}/2}} \prod_{a=1}^{N_{\text{train}}} \delta[y(\vec{\theta}_a) - E(\vec{\theta}_a)]$$

$$\prod_{i=1}^{N_{\text{coef}}} dA_i \ d\sigma_A^2 d\Lambda \exp\left\{-\frac{1}{2\sigma^2} A_i^2\right\}.$$
(9.19)

If the training constraints are not exact, the delta function should be replaced. This would be true if the training had some sort of point-by-point noise, such as one would have if the full-model calculation involved some sort of sampling or Monte Carlo evaluations. However, more generally one could also consider the fact that the training calculation might have some sort of error, e.g. that of missing physics. In that even the error might be correlated. If errors in the full-model calculations at points  $\vec{\theta}_a$  and  $\vec{\theta}_b$  are characterized some error matrix,

$$\Delta_{ab} = \langle \delta y_a \delta y_b \rangle, \tag{9.20}$$

Here,  $y_a = y(\vec{\theta}_a)$ , the full model value calculated at the training point a. The delta function above is replaced by

$$\prod_{a} \delta[y_a - E(\vec{\theta}_a)] \to \frac{|\Delta|^{1/2}}{(2\pi)^{N_{\text{train}}/2}} \exp\left\{\frac{-1}{2}(y_a - E(\vec{\theta}_a))\Delta_{ab}^{-1}(y_b - E(\vec{\theta}_b))\right\}$$

$$= \int \prod_{a} \left(\frac{dk_a}{2\pi}\right) \exp\left\{ik_a(y_a - A_iT_i(\vec{\theta}_a)) - \frac{1}{2}k_a\Delta_{ab}k_b\right\}.$$
(9.21)

Defining the net probability after integrating over the coefficients above, but not over  $\sigma_A^2$  or  $\Lambda$ ,

$$\mathcal{Z} \equiv \frac{dP(\sigma_A, \Lambda)}{d\sigma_A d\Lambda},\tag{9.22}$$

the average coefficients, and correlation of coefficients, for a given  $\sigma_A$  and  $\Lambda$ , can be expressed in the following form after replacing the delta function,

$$\langle A_{j} \rangle = \mathcal{Z}^{-1} \int \prod_{i} \frac{dA_{i}}{\sqrt{2\pi\sigma_{A}^{2}}} A_{j} \exp\left[-\sum_{\ell} \frac{A_{\ell}^{2}}{2\sigma_{A}^{2}}\right]$$

$$\int \prod_{a} \frac{dk_{a}}{2\pi} \exp\left[ik_{b}(y_{b} - A_{i}T_{i}(\vec{\theta}_{b})) - \frac{1}{2}k_{a}\Delta_{ab}k_{b}\right],$$

$$\langle A_{j}A_{k} \rangle = \mathcal{Z}^{-1} \int \prod_{i} \frac{dA_{i}}{\sqrt{2\pi\sigma_{A}^{2}}} A_{j}A_{k} \exp\left[-\frac{A_{i}^{2}}{2\sigma_{A}^{2}} - \frac{1}{2}k_{a}\Delta_{ab}k_{b}\right]$$

$$\int \prod_{a} \frac{dk_{a}}{2\pi} \exp\left[ik_{b}(y_{b} - A_{i}T_{i}(\vec{\theta}_{b}))\right],$$

$$\mathcal{Z} = \int \prod_{i} \frac{dA_{i}}{\sqrt{2\pi\sigma_{A}^{2}}} \exp\left[-\frac{A_{i}^{2}}{2\sigma_{A}^{2}}\right] \int \frac{dk}{2\pi} \exp\left[ik_{b}(y_{b} - A_{\ell}T_{\ell}(\vec{\theta}_{b})) - \frac{1}{2}k_{a}\Delta_{ab}k_{b}\right].$$

$$(9.23)$$

After completing the square, the correlation can be written as

$$\langle A_{j}A_{k}\rangle = \mathcal{Z}^{-1} \int \prod_{a} \frac{dk_{a}}{2\pi} e^{ik_{b}y_{b} - \frac{1}{2}k_{a}\Delta_{ab}k_{b}}$$

$$\int \prod_{i} dA_{i} A_{k}A_{j} \exp \left[ -\frac{1}{2\sigma_{A}^{2}} \left( A_{i} + i\sigma_{A}^{2}T_{i}(\vec{\theta}_{b})k_{b} \right)^{2} - \frac{\sigma_{A}^{2}}{2} \left( \sum_{\ell} T_{\ell}(\vec{\theta}_{b})k_{b} \right)^{2} \right]$$

$$= \mathcal{Z}^{-1} \int \prod_{a} \frac{dk_{a}}{2\pi} \exp \left[ -\frac{\sigma_{A}^{2}}{2} (T_{j}(\vec{\theta}_{b})k_{b})^{2} + ik_{b}y_{b} - \frac{1}{2}k_{a}\Delta_{ab}k_{b} \right]$$

$$\int \prod_{i} dA_{i} \left( A_{k} - i\sigma_{A}^{2}T_{k}(\vec{\theta}_{b})k_{b} \right) (A_{j} - i\sigma_{A}^{2}T_{j}(\vec{\theta}_{c})k_{c}) \exp \left[ -\sum_{\ell} \frac{A_{\ell}^{2}}{2\sigma_{A}^{2}} \right]$$

Similarly,

$$\langle A_j \rangle = -\mathcal{Z}^{-1} \int \prod_a \frac{dk_a}{2\pi} \exp\left[ -\frac{\sigma_A^2}{2} (T_j(\vec{\theta}_b) k_b)^2 + ik_b y_b - \frac{1}{2} k_a \Delta_{ab} k_b \right]$$

$$\int \prod_i \frac{dA_i}{\sqrt{2\pi\sigma_A^2}} i\sigma_A^2 T_j(\vec{\theta}_c) k_c \exp\left[ -\frac{A_\ell^2}{2\sigma_A^2} \right],$$

$$\mathcal{Z} = \int \prod_a \frac{dk_a}{2\pi} \exp\left[ -\frac{\sigma_A^2}{2} (T_j(\vec{\theta}_b) k_b)^2 + ik_b y_b - \frac{1}{2} k_a \Delta_{ab} k_b \right].$$
(9.25)

We define the operator,

$$B_{ab} \equiv \sum_{i} T_i(\vec{\theta}_a) T_i(\vec{\theta}_b) + \frac{\Delta_{ab}}{\sigma_A^2}.$$
 (9.26)

Doing the integrals for  $\mathcal{Z}$ ,

$$\mathcal{Z} = \sigma_A^{-N_{\text{train}}} (\det B)^{-1/2} \exp \left[ -\frac{1}{2\sigma_A^2} y_a B_{ab}^{-1} y_b \right]. \tag{9.27}$$

One can then find the moments of the integral over k,

$$\langle k_a \rangle = -i\partial_{y_a} \ln(\mathcal{Z}) = \frac{i}{\sigma_A^2} B_{ab}^{-1} y_b,$$

$$\langle \delta k_a \delta k_b \rangle = -\partial_{y_a} \partial_{y_b} \ln(\mathcal{Z}) = \frac{1}{\sigma_A^2} B_{ab}^{-1}.$$
(9.28)

The moments are then:

$$\langle A_j \rangle = T_j(\vec{\theta}_a) B_{ab}^{-1} y_b,$$

$$\langle \delta A_j \delta A_k \rangle = \delta_{jk} \sigma_A^2 - \sigma_A^2 B_{ab}^{-1} T_j(\vec{\theta}_a) T_k(\vec{\theta}_b).$$
(9.29)

The observables and their fluctuations are then:

$$\langle E(\vec{\theta}) \rangle = \langle A_j \rangle T_j(\vec{\theta}), \qquad (9.30)$$
$$\langle \delta E(\vec{\theta})^2 \rangle = \langle \delta A_j \delta A_k \rangle T_j(\vec{\theta}) T_k(\vec{\theta}).$$

After defining

$$\chi_a \equiv B_{ab}^{-1} y_b, \tag{9.31}$$

$$S_a(\vec{\theta}) \equiv T_j(\vec{\theta}) T_j(\vec{\theta}_a),$$

the mean and fluctuation can be written as

$$\langle E(\vec{\theta}) \rangle = \chi_a S_a(\vec{\theta}),$$

$$\langle \delta E(\vec{\theta})^2 \rangle = \sigma_A^2 T_j(\vec{\theta}) T_j(\vec{\theta}) - \sigma_A^2 S_a(\vec{\theta}) B_{ab}^{-1} S_b(\vec{\theta}).$$
(9.32)

Whereas  $\langle E(\vec{\theta}) \rangle$  depends linearly on the training values,  $y_{a=1,N_{\text{train}}}$ , the emulator's uncertainty,  $\langle \delta E(\vec{\theta})^2 \rangle$  depends only on the location of the training points.

If one sets  $\vec{\theta} = \vec{\theta}_c$ , one of the training points, and if  $\Delta = 0$ , then

$$S_a(\vec{\theta_c}) = B_{ac},$$
 (9.33)  
 $T_i(\vec{\theta_c})T_i(\vec{\theta_c}) = B_{cc}, \text{ no sum over } c,$ 

and

$$\langle \delta E(\vec{\theta_c})^2 \rangle = \sigma_A^2 B_{cc} - \sigma_A^2 B_{ca} B_{ab}^{-1} B_{bc} = 0.$$
 (9.34)

Thus, there is no uncertainty when evaluated at the training points. When  $\Delta \neq 0$ , the uncertainty at a training point,  $\vec{\theta_c}$  is

$$\langle \delta E(\vec{\theta_c})^2 \rangle = \Delta_{cc} - \frac{1}{\sigma_A^2} \Delta_{ca} B_{ab}^{-1} \Delta_{bc}, \quad \text{(no sum over } c\text{).}$$
 (9.35)

If  $\Delta$  is small, the uncertainty of the emulator is simple given by  $\Delta$ , however it is somewhat reduced for larger  $\Delta$ . Because the matrices  $T_i(\vec{\theta}_a)T_i(\vec{\theta}_b)$  and  $\Delta_{ab}$  are both positive definite, the uncertainty squared above is manifestly positive.

Before considering a specific position,  $\vec{\theta}$ , one first calculates and stores the vector  $\chi$  and the matrix  $B^{-1}$ . This involves inverting a  $N_{\text{train}} \times N_{\text{train}}$  matrix. Then, when considering a specific  $\vec{\theta}$ , one calculates  $S_a(\vec{\theta})$  for each a. This involves calculating a double sum with  $N_{\text{train}} \times N_{\text{coef}}$  elements. Once these are stored, calculating  $\langle \delta y^2 \rangle$  involves calculating a double sum with an additional  $N_{\text{train}} \times N_{\text{train}}$  elements. Thus, calculating  $S_a(\vec{\theta})$  is where the bulk of the calculation occurs. Because  $N_{\text{train}} \lesssim 100$  and  $N_{\text{coef}}$  might be of in the tens of thousands (depending on the number of parameters and maximum rank), one should be able to calculate  $\langle y \rangle$  and  $\langle \delta y^2 \rangle$  in much less than a second. Nonetheless, when performing an MCMC calculation, one might calculate for millions of values of  $\vec{\theta}$ , which can push the calculation into the ranges of minutes or even hours if one has a dozen parameters or more.

The calculations above could have been performed finding the coefficients to maximize the probability,  $P(\vec{A})$ , rather than the mean coefficients, but the answer is exactly the same.

#### 9.3.2 Choosing the Variance for the Coefficients and the Smoothness Parameter

Here, we find the optimum choice for  $\sigma_A$ , the variance of the coefficient in the absence of training data, in the previous section. For the derivations of  $\langle y \rangle$  and  $\langle \delta y^2 \rangle$  in the previous section, the expression in Eq. (9.29) for  $\langle A_i \rangle$  has no dependence on  $\sigma_A$ , which sets the variance of the coefficients. Further, the expression for  $\langle \delta A_i^2 \rangle$  does not depend on the model training values,  $y_{a=1,N_{\text{train}}}$ . The net probability from Eq. (9.36) was found to be in Eq. (9.19) to be

$$\int \prod_{i} dA_{i} P(\vec{A}, \sigma^{2}, \Lambda) = \mathcal{Z}$$

$$= \sigma_{A}^{-N_{\text{train}}} \left( \det B \right)^{1/2} \exp \left[ -\frac{1}{2\sigma_{A}^{2}} y_{a} B_{ab}^{-1} y_{b} \right].$$
(9.36)

Our goal here is to find the value of  $\sigma_A$  that maximizes  $\mathcal{Z}$ . For the case where  $\Delta = 0$ , one can find  $\sigma_A$  rather easily. In that case, setting the derivative w.r.t.  $\sigma_A^2$  equal to zero,

$$0 = \frac{N_{\text{train}}}{\sigma_A^2} \mathcal{Z} - \frac{\mathcal{Z}}{\sigma_A^4} y_a B_{ab}^{-1} y_b, \tag{9.37}$$

$$\sigma_A^2 = \frac{1}{N_{\text{train}}} y_a B_{ab}^{-1} y_b. \tag{9.38}$$

However, the situation is becomes more complicated if  $\Delta \neq 0$ . Fortunately, our tests show that the value of  $\sigma_A^2$  is only very slightly changed by random uncertainties if those uncertainties are less than 10 percent of  $\sigma_A$ . For that reason, *Smooth Emulator* ignores  $\Delta_{ab}$  while choosing  $\sigma_A$ . Tests (see last subsection) showed that the appropriate value of  $\sigma_A$  can usually be found to within 10%.

Choosing the smoothness parameter,  $\Lambda$ , is unfortunately not simple. Given the lack of an analytic solution, one can simply plot  $\ln(Z)$  vs  $\Lambda$ . However, it was found (as shown in the final subsection here) that afater using a function for the surrogate full-model characterized by a known  $\Lambda$ , the value at which  $\ln(Z)$  is minimized can vary significantly from the true value. Though this method reproduces the correct value if averaged over many possible surrogate functions, it varies too much on a per-emulation basis to be useful. For example, for some instances of sample functions, it was

found that  $\ln(Z)$  would be maximized for  $\Lambda < 1.5$ , even when the surrogate-model function was generated with  $\Lambda = 3$ .

Our conclusion (see final sub-section) regarding the ability  $Smooth\ Emulator$  to determine  $\sigma_A$  and  $\Lambda$  is that  $\sigma_A$  can be found quite robustly using the simple analytic form in Eq. (??), but that the determination of  $\Lambda$  is questionable. Fortunately, the predicted values of the emulated function are not strongly sensitive to  $\Lambda$ , but the estimated emulator uncertainties are somewhat sensitive. Thus, one can plot  $\ln(Z)$  vs  $\Lambda$  to get some insight, but the result should be taken with caution. It might make more sense to set  $\Lambda$  to some guess, perhaps 2.5, then just live with the knowledge that this might not be the best choice. If one can perform numerous additional full-model runs at points not used for training, one can choose  $\Lambda$  such that the uncertainties are large enough to capture 68% of the points within one standard deviation. From the experience of trying many functions, it would seem that optimizing  $\Lambda$  in this way can improve the uncertainty estimates by approximately 50%. I.e. changing  $\Lambda$  by factors of 2 or 4, tends to change the estimated uncertainties by tens of percent.

# 10 Theoretical Basis of Simplex Sampler

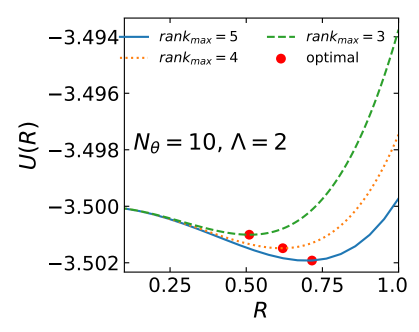
Here, we imagine a spherically symmetric prior, e.g. one that is purely Gaussian, and where the parameters are scaled so that the prior distribution is invariant to rotations. If one believe the function were close to linear, a strategy would be to find  $N_{\text{train}} = N + 1$  points in parameter space placed far apart from another. One choice is the N-dimensional simplex. Examples are an equilateral triangle in two-dimensions or a tetrahedron in three dimensions. For a simplex, one places the  $N_{\text{train}} = N + 1$  training points at a uniform distance from the origin, r, with equal separation between each point. One can generate an N-dimensional simplex from an N-1 dimensional arrangement. In the N-1 dimensional arrangement, the points are arranged equidistant from one another using the coordinates  $x_1 \cdots x_N$ . The points would all be placed at a radius  $r_N$  from the origin in this system, and the separation would be d. In the N-dimensional system all these N-1 points had coordinate  $x_N = -a$ . The  $(N+1)^{\text{th}}$  point is then placed at position  $x_1 \cdots x_N = 0$  and  $x_{N+1} = Na$ . This keeps the center of mass at zero. One then chooses a such that the new point is equally separated from all the other points by the same distance d.

$$d^{2} = r_{N-1}^{2} + N^{2}a^{2},$$

$$a = \sqrt{\frac{d^{2}}{N^{2}} - r_{N-1}^{2}}.$$
(10.1)

Now, each of the N points is located a distance Na away from the center of the N-dimensional origin. This procedure can applied iteratively to generate the vertices of the simplex.

One might also wish to use enough training points to uniquely determine the emulator in the case that the function is quadratic. There are N(N+1)/2 additional points, which is exactly the number of segments connecting the N+1 equally-spaced vertices of the N-dimensional simplex. If placed at the midpoints of the segments, these points would be closer to the origin than the vertices. One of the simplex options is to place these points at the midpoints, then expand their radii while maintaining their direction. This would result in arrays of points at two different radii, with N+1 points positioned at the lower radius and N(N+1)/2 points being placed at the larger radius.



**Figure 10.1:** The integrated uncertainty for training with points set in a simplex geometry is found to depend only weakly on the radius. A Gaussian prior was assumed. The weak dependence was used to justify using a rather ad-hoc choice of radii for the simplex points, or as in the case of type 2, a second radius for the bisecting points.

From symmetry considerations, this method should produce either an extrema of effectiveness in regards to angular positioning (assuming Gaussian priors, which are rotationally invariant). I.e. any repositioning of a training point by a small differential  $d\vec{\theta}$  should not change the effectiveness if that differential does not change the radius of the point. However, one can expand all the radii of all the inner-positioned simplex, or all the radii of the other points, without changing the symmetry. Thus, the radii must be chosen to maximize the constraining power of the training data. To that end, a study was pursued where the overall integrated uncertainty was minimized as a function of the radius. Only the case of the simple simplex was considered, and only for the case of Gaussian priors. The minimum was found to depend on the maximum rank of the emulator, the smoothness parameter, and the number of observables. However, the minimum was always very broad, as shown in Fig. 10.1. For that reason, the radii were chosen rather ad hoc:

- Simple Simplex (N + 1 points, Type 1): The radii of each point was chosen to be 1 - 1/(N + 1).
- Simplex plus bisecting points (N+1) inner points plus N(N+1) outer points: The outer points were positioned at a radius 1-1/(N+1), while the inner points were positioned at a radius which was smaller by a factor of  $1/\sqrt{2}$ .

One issue with the simplex is that the first set of N+1 training points would all be placed at the same radius. If the prior parameter distribution is uniform within an N-dimensional hyper cube, the training points could be rather far from the corners in that space. Issues with such priors are discussed in the next section.

## 10.1 The Pernicious Nature of Step-Function Priors in High Dimension

For purely Gaussian priors, one can scale the prior parameter space to be spherically symmetric. Unfortunately, that is not true for step function priors (uniform within some range). In that case the best one can do (if the priors for each parameter are independent) is to scale the parameter space such that each parameter has the constraint,  $-1 < \theta_i < 1$ . If the number of parameters is N, the hyper-cube has 2N faces and  $2^N$  corners, a face being defined as one parameter being  $\pm 1$  while the others are zero, while a corner has each parameter either  $\pm 1$ . For 10 parameters, there are 1024 corners, and for 15 parameters there are 32678 corners. Thus, it might be untenable to place a training point in each corner.

One can also see the problem with placing the training points in a spherically symmetric fashion as is done with the *Simplex Sampler*. The hyper-volume of the parameter-space hyper-cube is  $2^N$ , whereas the volume of an N-dimensional hyper-sphere of radius R = 1 is

$$V_{\text{sphere}} = \Omega_N \int_0^R dr \ r^{N-1} = \Omega_N \frac{R^N}{N}. \tag{10.2}$$

The solid angle,  $\Omega_N$  in N dimensions is

$$\Omega_N = \frac{2\pi^{N/2}}{\Gamma(N/2)},\tag{10.3}$$

and after putting this together, the fraction of the hyper-cube's volume that is within the hyper-sphere is

$$\frac{V_{\text{sphere}}}{V_{\text{cube}}} = \begin{cases} \frac{(\pi/2)^{N/2}}{N!!}, & N = 2, 4, 6, 8 \dots \\ \frac{(\pi/2)^{(N-1)/2}}{N!!}, & N = 3, 5, 7, \dots \end{cases}$$
(10.4)

In two dimensions, the ratio is  $\pi/4$ , and in three dimensions it is  $\pi/6$ . In 10 dimensions it is  $2.5 \times 10^{-3}$ . For high dimensions only a small fraction of the parameter space can ever lie inside inside a sphere used to place points. And, if the model is expensive, it may not be tenable to run the full model inside every corner.

One can also appreciate the scope of the problem by considering the radius of the corners vs. the radius of the sphere. The maximum value of  $|\vec{\theta}|$  is  $\sqrt{N}$ . So, for 9 parameters, if the training points were all located at positions  $|\vec{\theta}| < 1$ , one would have to extrapolate all the way to  $|\vec{\theta}| = 3$ . Thus, unless the model is exceptionally smooth, one needs to devise a strategy to isolate the portion of likely parameter space using some original set of full-model runs, then augment those runs in the likely region.

A third handle for viewing the issue in N dimensions is to compare the r.m.s. radii of the hypersphere to that of the hyper-cube. For the cube where each side has length 2a,

$$\left(R_{\text{r.m.s.}}^{\text{(cube)}}\right)^2 = a^2 \frac{N}{3}.\tag{10.5}$$

whereas for a sphere of radius a,

$$\left(R_{\text{r.m.s.}}^{\text{(sphere)}}\right)^2 = a^2 \frac{N+2}{N}.$$
(10.6)

The ratio of the radii is then

$$\frac{R_{\text{r.m.s.}}^{\text{(sphere)}}}{R_{\text{r.m.s.}}^{\text{(cube)}}} = \sqrt{\frac{3}{N+2}}.$$
(10.7)

Thus, in 10 dimensions, if the training points are placed at a distance a from the origin, the r.m.s. radii of the interior space would be half that of the entire space. Further, the r.m.s. radii of the points in the cube,  $a\sqrt{N/3}$ , would be about 83% higher than the radius of the training points.

Of course, these problems would be largely avoided if the number of parameters was a half dozen or fewer, or if one was confident that the function was extremely smooth. In the first two sections of this paper, the smoothness parameter,  $\Lambda$ , was set to a constant. There might be prior knowledge that certain parameters affect the observables only weakly. In that case, the response to these parameters can be considered as linear. This could be done by scaling those parameters so that they vary over a smaller range. If a parameter varies only between -0.1 and 0.1, that effectively applies a smoothness parameter in those directions that is ten times higher. Unfortunately, the choice of which parameters to rescale in this fashion would likely vary depending on which observable is being emulated. Because all the observables might be calculated in a full model run, one needs to identify parameters that would likely have weak response on all observables.

## 11 Tests of Smooth Emulator

Here, we show several results for testing  $Smooth\ Emulator$ . For these tests we consider a surrogate full-model generated from Taylor expansions with randomized coefficients, exactly according to the form assumed by the emulator. These functions are characterized by a smoothness parameter,  $\Lambda$ , a variance for the coefficients,  $\sigma_A$ , and a maximum rank. If the emulator works correctly, 68% of the discrepancies in emulated functions, compared to the true model, should fall within one standard deviation. Performing the same test as described in the tutorial, Fig. 11.1 compares emulator predictions to 50 different surrogate models. Each surrogate model was constructed according to Taylor expansions with  $\Lambda = 3$ , a maximum rank of 4, and  $\sigma_A = 100$ . Although only 50 such models are illustrated here, this was attempted with 5000 models, and it was found that the percentage of predictions within one sigma was within a few tenths of a percent of the 68% expected, thus validating both the emulator's prediction of both its value and its estimate of its uncertainty.

Figure 11.2 shows the average of surrogate-model predictions  $\ln(Z)$ ,  $\sigma_A$  and the percentage of predictions within one sigma over 500 instances. These are shown as a function of the emulator's  $\Lambda$ . Because the surrogate models were constructed with  $\Lambda = 3$ , the value of  $\ln(Z)$  should be a maximum there. Further, for  $\Lambda = 3$ , the extracted value of  $\sigma_A$  should be 100, and 68% of emulator predictions should be within the emulator's uncertainty estimate. The emulator passes all these tests beautifully, but as can be seen, whereas the prediction of  $\sigma_A$  is remarkably accurate, the value of  $\ln(Z)$  is quite uncertain. Thus, finding  $\Lambda$  for a given emulator might be problematic. Figure 11.3 shows  $\ln(Z)$  for several individual models. This was repeated for 5, 10 and 15 parameters. For smaller numbers of parameters, using  $\ln(Z)$  vs  $\Lambda$  to determine  $\Lambda$  is especially problematic. Thus, the User might want to simply fix  $\Lambda$  at some value, e.g 2.5, and live with the consequences.

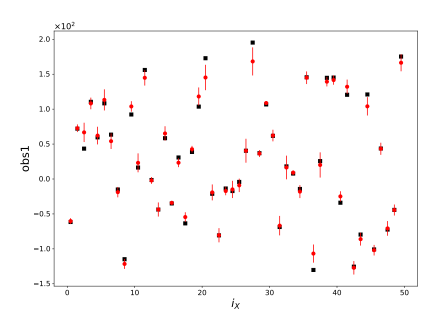
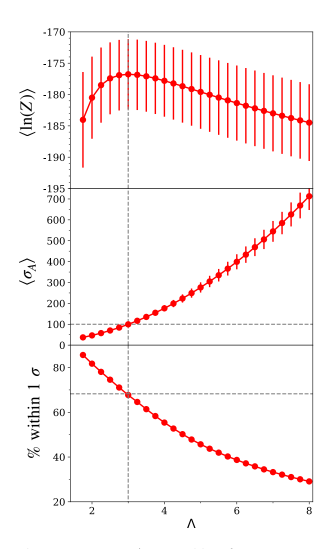


Figure 11.1: Emulator predictions are compared to a surrogate model with Taylor expansions based on  $\Lambda = 3$  and  $\sigma_A = 100$ . As can be seen approximately 68% of the predictions were within one sigma of the true model, validating the emulator, both in regards to its predicted value and its uncertainty estimate.



 $N_{par} = 5$ -60 -70 -80 -90 -100 -110 <u>N</u> -120 -130  $I_{\rm par} = 15$ -140 -150 -160 -170 -180  $\dot{4}$ 6 8 ٨

Figure 11.2: A ensemble of 5000 surrogate models, each generated with  $\Lambda = 3$  an  $\sigma_A = 100$  were emulated. The extracted values of  $\ln(Z)$  an  $\sigma_A$ , along with the percentage of emulator predictions withing the emulator's uncertainty were plotted agains the value of  $\Lambda$  assumed for the emulator. Indeed, for  $\Lambda = 3$ ,  $\ln(Z)$  is at a maximum,  $\sigma_A = 100$  and 68% of the predictions are within the estimated uncertainty. Unfortunately,  $\ln(Z)$  has a high instance-to-instance variability.

Figure 11.3: For 20 instances of the surrogate function,  $\ln(Z)$  is plotted vs.  $\Lambda$ . Although if one were to average over thousands of surrogate models, one could extract  $\Lambda$  accurately, the extraction is rather inaccurate on a model-by-model basis. This is especially true if the model has smaller numbers of parameters.