Building Emulators

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https://github.com/andrewcparnell/intro_emulators

Introduction

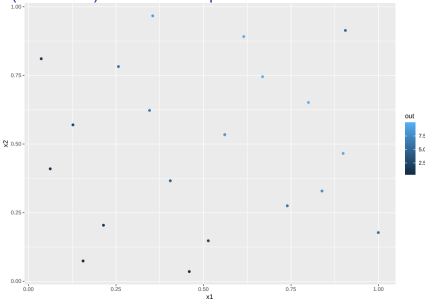
- ▶ We have now chosen our best input values to run the simulator at
- ▶ We now need to run the simulator. This may take a while...
- ▶ ... but then we can build our emulator!
- We will use a Gaussian Process to build the emulator

First step: run the simulator

- Assume we have chosen our input points using a suitable design and that we have chosen our design points
- ▶ We now run our simulator for sine wave example with:

```
n_runs <- 20
n_inputs <- 2
initial_grid <- maximinLHS(n_runs, n_inputs)
# Set up container and run
out <- rep(NA, length = n_runs)
for (i in 1:n_runs) {
   out[i] <- f(initial_grid[i, 1], initial_grid[i, 2])
}</pre>
```

Example 1 (sine wave) simulator output



Example 2 (2D Navier-Stokes) simulator run

```
for (i in 1:n runs) {
  system("./dockerClean2.sh")
  curr answer <- system(paste("./dockerRun.sh",</pre>
                          df_grid$largeInletVelocityMagnitude[i],
                          df grid$smallInletVelocityMagnitude[i],
                          df_grid$kinematicViscosity[i]),
                    intern = TRUE
  df_grid$out[i] <- as.numeric(curr_answer[length(curr answer) - 1])</pre>
  print(df grid$out[i])
```

Fitting the emulator

- \blacktriangleright We now have a set of outputs (y) and a set of inputs (X)
- ▶ We assume as before that *y* is univariate and *X* is multivariate
- ► This is now a standard machine learning problem! (Go and see the Rfternoon series of introductory R lectures course if you want to solve machine learning problems)
- ... but mostly these will not work!
- This is because the simulator is deterministic we want our emulator to reproduce f(X) = y when we choose values of X that we have already run
- ▶ Most machine learning algorithms do not do this, except for Gaussian Processes. . .

What is a Gaussian Process?

- ► A Gaussian Process is just a fancy version of linear regression where we also model the correlation between the response values using a **multivariate normal distribution**
- ► This is the model:

$$y|x \sim MVN(\mu 1, \Sigma); \ \Sigma_{ij} = \tau^2 \rho_{\phi}(\|x_i - x_j\|)$$

- ▶ There are 3 parameters: μ, ϕ, τ
- ▶ The parameter τ^2 controls the variance of the Gaussian process (bigger values have more uncertainty)
- ightharpoonup The parameter ϕ controls how smooth the curve is
- The function ρ controls the correlation function; how the correlation between the y values decreases as x gets further away

Gaussian process predictions

▶ The GP has a wonderful formula for predicting at new input values x^* :

$$\hat{y}^*|y \sim \textit{MVN}(\mu 1 + \Sigma^* \Sigma^{-1}(y - \mu 1), \Sigma^{**} - \Sigma^* \Sigma^{-1}(\Sigma^*)^T)$$

- ▶ Here Σ^* and Σ^{**} are the covariance matrix of the prediction points and their variance respectively
- If you are predicting for a known point then $\Sigma^{**} = \Sigma^* = \Sigma$ and the prediction is just y. This means it will go through the points you have already run in your simulator!
- ► The other advantage is that you get predicted means and uncertainties from every point you choose

Why Gaussian processes?

- ► There is nothing special about the fact that it's a Gaussian Process used here. Any statistical model that can produce predictions with quantified uncertainties should be OK, provided it produces zero variance predictions at given input values
- ► Another nice thing about GPs is that they are smooth which often matches the differential equations underlying the simulator
- ► The bad thing about them is that they can be slow to fit we need to solve the covariance matrix when optimising the parameters and creating predictions

The GPFit package

- ▶ We will use the GPfit package to create our Gaussian Process predictions because it's very simple
- Go back and see part 1 for richer alternatives
- ▶ One downside to using GPfit is that it requires all inputs to be scaled between 0 and 1. This just requires an extra few lines of coding

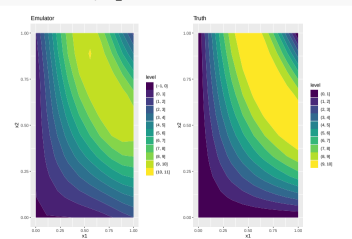
Fitting

Fitting the model is just one line!

```
emulator <- GP_fit(initial_grid, out)</pre>
```

For a 2 input example we can plot the emulator, but we need to create our own plots after that

Predicting new values for example 1



Emulating via other machine learning models

- ➤ You don't have to use a GP to fit your emulator, you could use your favourite machine learning algorithm
- ▶ But remember that it will likely not give the true simulator values at the same input
- ▶ Using a different machine learning approach will work well if (a) you have a stochastic simulator, or (b) you can perform a very large number of simulation runs

Summary

- ▶ We have now fitted a Gaussian Process emulator to our example data
- Gaussian processes are just a clever machine learning tool that are both smooth and exhibit zero variance at the input values
- Using the GPfit package it is only one line to fit an emulator
- ► Can then go and predict for any new values of the simulator more or less instantly
- Next: plotting, checking, and calibrating the emulator...