Overview

- Metamodels
- Gaussian Process background
- Gaussian Process formulation
- Examples
- Use of Gaussian Processes within other methods (e.g. optimization)
Metamodels

- Metamodels: also called surrogate, response surface model, or emulator.

- Typically constructed over a small number of simulation model runs (“code runs”)

- The simulation is very costly to run, we can only afford a limited number of runs, often dozens to a few hundred

- The code runs provide the training data (e.g. sets of input parameters and corresponding response values)

- The metamodel is constructed to provide a fast, cheap function evaluation for the purposes of uncertainty quantification, sensitivity analysis, and optimization.

Metamodels

- Taylor series approximations
- Linear regression models
- Neural networks
- Moving least squares
- Radial Basis Functions
- Multivariate Adaptive Regression Splines (MARS)
- Gaussian process models
- Polynomial Chaos expansions
- Multi-fidelity models
- Reduced-order models
Gaussian Processes

- Why are GPs popular emulators of computer models?
  - They allow modeling of fairly complicated functional forms
  - They do not just offer a prediction at a new point but an estimate of the uncertainty in that prediction

- Classic references:
    - http://www.gaussianprocess.org/gpml/chapters/
Gaussian Process

- A stochastic process is a collection of random variables \( \{y(x) \mid x \in X\} \) indexed by a set \( X \) in \( \mathbb{R}^d \), where \( d \) is the number of inputs.

- A Gaussian process is a stochastic process for which any finite set of \( y \)-variables has a joint multivariate Gaussian distribution. That is, the joint probability distribution for every finite subset of variables \( y(x_1), \ldots, y(x_k) \) is multi-variate normal.

- A GP is fully specified by its mean function \( \mu(x) = E[y(x)] \) and its covariance function \( C(x, x') \).
What does this mean?

- Start with a set of runs of a computer code: at each sample $x_i$ we have output $y_i(x_i)$.
- The output at a new input value, $x_{\text{new}}$, is uncertain.
- This is what a GP will predict.
- Related to regression.
- Related to random functions. From our set of samples, we have a “deterministic” function that is a set of points $\{x, y(x)\}$ or $\{x, f(x)\}$. Instead of $f(x)$, if we use the outcome of a random draw from some joint distribution of random variables $\{Z(x_1), \ldots, Z(x_n)\}$, we get a realization of a random function.
- This is a stochastic process (e.g. generate many draws and get many functions).
How do we simulate realizations of a random function?

- Start with \( \{Z(x_1), \ldots, Z(x_n)\} \) from a multivariate normal distribution with mean 0 and covariance matrix \( C = \text{Cov}[Z(x_i), Z(x_j)] \).

- To simulate a random draw:
  - Generate \( n \) standard normal(0,1) random variables, \( S \).
  - Perform a Cholesky decomposition \( C = LL^t \).
  - Define \( Z = LS \).
  - Plot the points \( \{x_i, Z_i = Z(x_i)\} \)
  - Connect the dots
Example covariance function in 1-D

- \( \text{Cov}[Z(x_i), Z(x_j)] = \exp(-\theta |x_i - x_j|^2) \)
Gaussian Process

- We have the capability to generate random functions
- We can add a mean function (typically a constant or a simple polynomial regression)
- We can multiply the covariance by a constant to scale the vertical axis.
- Now, we can vary $\theta$ to get a certain amount of “wiggle” in the random function (smaller $\theta$ leads to less wiggle).
- **NOW:** we want to constrain these random functions to be consistent with the data points we have
- We can either take a Bayesian approach or a maximum likelihood (MLE) approach to estimate the parameters governing the Gaussian process
- Start with a MLE approach
Gaussian Process

- Typical formulation: a Gaussian process is defined by its mean and covariance function. We assume:
  \[ E[y(x)] = f(x)^T \beta \quad \text{Mean} \]
  \[ \text{Cov}[y(x), y(x')] = \sigma^2 r(x, x') \quad \text{Covariance} \]
  \[ Y \sim N(f(X)^T \beta, \sigma^2 R) \quad \text{Multivariate Normal} \]

- A few notes:
  - \( x \) is one set of inputs of dimension \( d \). We have \( N \) samples, \( x_i \), for \( i=1...N \). Each \( x_i = \{x_{i1}, x_{i2}, ..., x_{id}\} \). \( X \) denotes the \((d*N)\) set of all samples, and \( \beta \) is the \( d*1 \) vector of regression coefficients. It may just be a constant \( \beta \).
  - It is more typical to write the covariance as the product of a scaling factor \( \sigma^2 \) times the correlation \( r(x, x') \).
  - The full \( N \times N \) correlation matrix between all points is \( R \).
  - \( Y \) is the \((N*1)\) vector of response values.
Gaussian Process

- NOW: what is the prediction for a new point?

\[ E[y(x^*)|Y] = f(x^*)^T \beta + r(x^*)^T R^{-1} [Y - F \beta] \]
\[ \text{Var}[y(x^*)|Y] = \sigma^2 (1 - r(x^*)^T R^{-1} r(x^*)) \]

- The correlation matrix for the training points is \( R \).
- \( r(x^*) \) is the vector of correlations between the new point \( x^* \) and the existing N points. It is of size N*1.
- \( F \) is the set of basis functions for the original full data set \( X \).
- These are the **conditional predictions** (conditional on the data).
What does this look like?

Note the reduction in variance as you have more data.
What does this look like?

This plot shows mean and variance plus random realizations.

https://pythonhosted.org/infpy/gps.html
Properties of the GP approximation

- The mean prediction interpolates the data.
  \[ E[y(x^*)|Y] = f(x^*)^T \beta + r(x^*)^T R^{-1}[Y - F\beta] \]

- The mean prediction is a linear combination of basis functions

- The predicted variance increases the further away the new point is from existing points.
  \[ \text{Var}[y(x^*)|Y] = \sigma^2 (1 - r(x^*)^T R^{-1} r(x^*)) \]
Correlation Function

- Want to capture the idea that nearby inputs have highly correlated outputs.
- The correlation in some dimensions may be more important than others...different “length-scales” in each dimension
- Common correlation functions include

**Power-exponential (or squared exponential):**

- Typically the exponent $p_j$ is 2, which gives smooth realizations. If $p_j$ is 1, you get much rougher realizations.
- Larger values of $\theta_j$ mean smaller correlation in the $x_j$ direction.

$$R(x, x') = \exp\{-\sum_{j=1}^{d} \theta_j (x_j - x_j')^{p_j}\} = \prod_{j=1}^{d} \exp(-\theta_j (x_j - x_j')^{p_j})$$
Correlation Function

Matern

\[ R(x, x') = \prod_{j=1}^{d} \frac{2^{1-\nu}}{\Gamma(\nu)} (-\theta_j |x_j - x_j'|^{\nu}) K_{\nu} (-\theta_j |x_j - x_j'|^{\nu}) \]

- Is equal to the exponential covariance function when \( \nu = \frac{1}{2} \).
- Is equal to the squared exponential when \( \nu \to \infty \).
- Typically, \( \nu = \frac{1}{2}, 3/2, \) or \( 5/2 \), going a process that looks rough to a process that is fairly smooth.
- Other covariances are possible: Cauchy, polynomial functions, etc.
Putting it all together

- Start with $N$ runs of a computer code, with points $\{x_i, y_i\}$. Ideally, the $N$ points will be a well-spaced design such as Latin Hypercube.

- Define the mean function for the Gaussian process.
  - Often, zero mean or constant mean is used.

- Define the covariance function for the Gaussian process.
  - Typically, the power-exponential function is used.

- Estimate the parameters governing the Gaussian process, including $\beta$, $\sigma$, and any parameters of the correlation function $R$ such as $\theta_j$.
  - Can use maximum likelihood or Bayesian methods

- Substitute the parameters in the prediction equations and obtain mean and variance estimates for new points $x^*$
Parameter Estimation (MLE)

- The observed training values represent a realization of a multivariate normal distribution.

\[ f(Y) = (2\pi)^{-\frac{N}{2}} |\Sigma|^{-\frac{1}{2}} \exp \left( -\frac{1}{2} (Y - \mu)^T \Sigma^{-1} (Y - \mu) \right) \]

- The basic idea of MLE is to find the particular mean vector and covariance matrix that define the most likely multivariate normal distribution to result in the observed data.

- Take the Log Likelihood and maximize it:

\[ \log(f(Y)) = -\frac{N}{2} \log(2\pi) - \frac{1}{2} \left( \sigma^2 N |R| \right) - \frac{1}{2\sigma^2} (Y - F\beta)^T R^{-1} (Y - F\beta) \]

- Drop the -1/2 term, and the first constant term and minimize the negative log-likelihood:

\[ NLL = N \log(\sigma^2) + \log(|R|) + \frac{1}{\sigma^2} (Y - F\beta)^T R^{-1} (Y - F\beta) \]
Parameter Estimation (MLE)

- Use global optimization methods to optimize the NLL OR
- Use gradient-based optimization to optimize the NLL. The derivations have been worked out with respect to $\beta$, $\sigma$, and correlation parameters of $R$.
- Conditional on fixed values of the correlation parameters, the optimal values for $\beta$ and $\sigma$ are given by the generalized least squares formulation:

$$\hat{\beta} = (F^T R^{-1} F)^{-1} (F^T R^{-1} Y)$$
$$\hat{\sigma}^2 = \frac{1}{N} (Y - F\beta)^T R^{-1} (Y - F\beta)$$

- One can use an iterative method, and obtain optimal correlation parameters $\theta$, then calculate $R$ and substitute it into above expressions above for $\beta$ and $\sigma$.
Bayesian parameter estimation

- Denote all of the parameters governing the GP as:
  - $\Theta = (\beta, \sigma, \theta_j)$.
- Bayesian approach to estimate posterior distribution on hyperparameters $\Theta$:
  \[
  \pi(\Theta \mid X, Y) \propto \pi(\Theta)L(X, Y \mid \Theta)
  \]
- Likelihood is the same as before with MLE
- Use Markov Chain Monte Carlo (MCMC) to solve it
- Requires thousands of evaluations of the likelihood function
- Large amount of work done in the statistical community about priors on these parameters, estimation of marginal likelihoods.
- Jeffreys-independent prior, reference priors, are often assumed
- Need to be careful that priors are not improper
Experimental Design

- The training set of \( \{x_i\} \) points, \( i = 1 \ldots N \) is usually a space-filling design such as a Latin Hypercube design or a maxi-min LHS.
- Want the points to be well spaced.
- Don’t want highly collinear points (close together).
- **PROBLEM:**
  - The prediction calculations require the inversion of the correlation matrix.
  - Often the correlation matrix is ill-conditioned and may be numerically singular.
  - Happens even with a few hundred points in 2-D.
  - One can’t invert \( R \) to use in the prediction calculation.
Techniques to handle ill-conditioning of the correlation matrix

- Remove points in a random or structured way ("Sparsification")
- Often, a small "jitter" or noise term $\sigma_\varepsilon$ is added to the diagonal terms of the covariance matrix to make the matrix better conditioned.

$$C = \sigma^2 R \rightarrow C = \sigma^2 R + \sigma_\varepsilon^2 I,$$

- Adding a nugget term
  - Estimate the nugget as part of the measurement error
  - Fix the measurement error and **add a nugget**, may have to do this iteratively until the nugget is big enough to make R well-conditioned
Techniques to handle ill-conditioning of the correlation matrix (cont’d)

- Linear algebra tricks
  - Don’t take the inverse of $R$, take the Cholesky factorization
  - Pseudo-inverse
    - Discards small singular values
  - Pivoted Cholesky Factorization
    - discard additional copies of the information that is most duplicated
    - Decrease the maximum eigenvalue and increase the minimum eigenvalue
  - Gradient-enhanced kriging

Software and Resources
Resources

- **Websites:** www.gaussianprocess.org
- **Managing Uncertainty in Computer Models (MUCM):**
  - UK project headed by Prof. Tony O’Hagan, University of Sheffield
  - http://www.mucm.ac.uk/Pages/ReadingList.html
- **Books:**
Software and Resources

- **Software:**
  - R:  `tgp` (Gramacy and Lee), `gptk` (Kalaitzis, Lawrence, et al.), `GPfit` (MacDonald, Chipman, and Ranjan)
  - Matlab: `gpml` (Rasmussen, Williams, Nickisch), `GPmat` (Sheffield Group)
  - Python: `GPy`, `gptools`, `pyGPs`, etc.
  - C++: [https://github.com/mblum/libgp](https://github.com/mblum/libgp)
  - MIT Group: MUQ/GPEXP (Python)
  - Dakota/Surfpack (C++)
  - Lots of others....
Example Use Cases
Efficient Global Optimization

- Technique due to Jones, Schonlau, Welch
- Build global Gaussian process approximation to initial sample
- Balance global exploration (add points with high predicted variance) with local optimality (promising minima) via an “expected improvement function”

\[ E[I(x)] = E\left[ \max(f_{\text{min}} - Y, 0) \right] \]

\[ E[I(x)] = (f_{\text{min}} - \hat{Y}) \Phi \left( \frac{f_{\text{min}} - \hat{Y}}{s} \right) + s \phi \left( \frac{f_{\text{min}} - \hat{Y}}{s} \right) \]

- Iteratively add points that have maximized EI, we use a DIRECT global optimization algorithm to identify that point
- Derivative-free, very efficient for low-dim.
Efficient Global Reliability Analysis (EGRA)

- Reliability methods find “failure surface” or “limit state contour” between “safe” and “failure” regions, often defined as $g(x) = 0$
- Integral of the probability density of the inputs over the failure region is the probability of failure

$$p_f = \int \cdots \int f_X(x_1, x_2, \ldots, x_n) \, dx_1 \, dx_2 \ldots dx_n$$

- Local reliability methods have problems with the nonsmooth, multimodal, and highly nonlinear failure surfaces
- EGRA is a global reliability analysis that uses a variant of EGO
  - The expected improvement is now the expected feasibility: penalize points from being away from the $g(x) = 0$ boundary
  - Balance explore and exploit in locating the limit state (EGRA)
  - Handles nonsmooth, multi-modal, highly nonlinear response functions
Efficient Global Reliability Analysis

Gaussian process model of reliability limit state with
10 samples

exploit

explore

28 samples
Efficient Global Reliability Analysis

Mean

Variance

Expected Feasibility
EGRA: Benchmark performance

Accuracy similar to exhaustive sampling at cost similar to local reliability assessment
Bayesian Formulation

- Generate posterior distributions on model parameters, given
  - Experimental data
  - A prior distribution on model parameters
  - A presumed probabilistic relationship between experimental data and model output that can be defined by a likelihood function

\[ \pi(\theta \mid d) \propto \pi(\theta) L(d \mid \theta) \]
Bayesian Calibration of Computer Models

- Experimental data = Model output + error

\[ d_i = M(\theta, x_i) + \epsilon_i \]

- If we assume error terms are independent, zero mean Gaussian random variables with variance \( \sigma^2 \), the likelihood is:

\[
L(\theta) = \prod_{i=1}^{n} \frac{1}{\sigma \sqrt{2\pi}} \exp \left[ - \frac{(d_i - M(\theta, x_i))^2}{2\sigma^2} \right]
\]

- How do we obtain the posterior?
  - It is usually too difficult to calculate analytically
  - We use a technique called Markov Chain Monte Carlo (MCMC)
  - In MCMC, the idea is to generate a sampling density that is approximately equal to the posterior. We want the sampling density to be the stationary distribution of a Markov chain.
Bayesian Calibration: Approach

- Take initial set of samples from simulation
  - Use LHS or space-filling design
- Develop Gaussian process approximation of the simulation
- Put priors on the input parameters
- Perform Bayesian analysis using MCMC
- Generate and analyze posterior distributions
- NOTE: GP surrogate adds a layer of uncertainty. However, this is explicitly modeled in the revised likelihood:

\[
L(\Theta) = 2\pi^{-n/2} \left| \Sigma \right|^{-1/2} \exp \left[ -\frac{1}{2} (d_i - \mu_{GP})^T \Sigma^{-1} (d_i - \mu_{GP}) \right]
\]

\[
\Sigma = \sigma^2 I + \Sigma_{GP}
\]

- Total uncertainty = (observation + model uncertainty) + surrogate uncertainty
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THANK YOU!

QUESTIONS?