# A tutorial introduction to the design and analysis of computer experiments

Darren Wilkinson darrenjw.github.io

# Computer models

## Computer experiments

- Due to the expense of running complex codes, it is necessary to carefully design experiments for computer models, somewhat analogous to physical designed experiments
- Just as we perform real experiments to reduce our uncertainty about the behaviour of physical systems, we can perform experiments on computer models to reduce our uncertainty about their (pre-determined, but unknown) behaviour
- Computer models often have various input parameters, which we denote by x, which alter the behaviour of the computer model
- Models will often generate lots of output. For this session, we
  will consider a single output of primary interest, y, which is a
  deterministic function of the inputs (even if there is some
  stochasticity internal to the computer model)

## **Emulating a computer model**

- An emulator is a cheap (fast) surrogate for the computer model, which has many potential uses
- Principally, it can be used instead of the real model in algorithms that require many evaluations of the simulator at many different input configurations
- Some emulators are good and some are bad to be useful, they should have a measure of uncertainty associated with them
- Stochastic process emulators define a probability distribution on the space of functions consistent with any known evaluations of the simulator
- Gaussian processes (GPs) are one of the simplest and most tractable stochastic processes on function space

# Gaussian processes (GPs)

## Gaussian processes

- A GP is a probability distribution on functions defined so that the marginal distribution of any finite number of points always has a multivariate normal (MVN) distribution
- Points close together in input space are typically more highly correlated than points far away
- Stationary GPs are defined by a covariance function many different possible choices - here I've used a Gaussian kernel

$$Cov[f(\mathbf{x}), f(\mathbf{x}')] = K(\mathbf{x}, \mathbf{x}') = \sigma^2 \exp \left\{ -\left(\frac{\|\mathbf{x} - \mathbf{x}'\|_2}{r}\right)^2 \right\},\,$$

containing two parameters: an asymptotic variance,  $\sigma^2$ , and a correlation length scale, r.

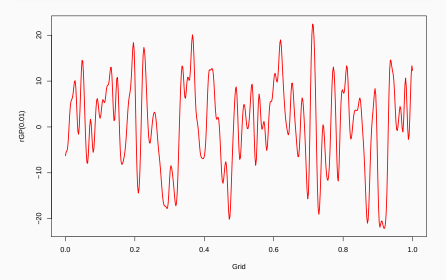
• A GP conditioned on observations is also a GP (Kriging)

#### 1d GP

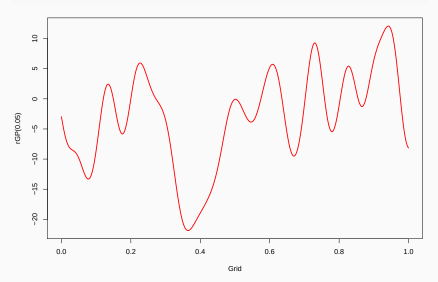
1d GPs are simplest to visualise and understand

```
Grid = seq(0,1,0.002)
Dist = as.matrix(dist(Grid))
K = function(d, lengthScale=0.4, scale=10)
  (scale^2)*exp(-(d/lengthScale)^2)
rGP = function(r=0.4, s=10)  {
  Var = K(Dist,r,s) + (1e-5)*diag(length(Grid))
  t(chol(Var)) %*% rnorm(length(Grid))
set.seed(1)
```

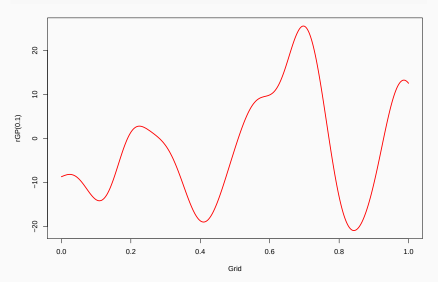




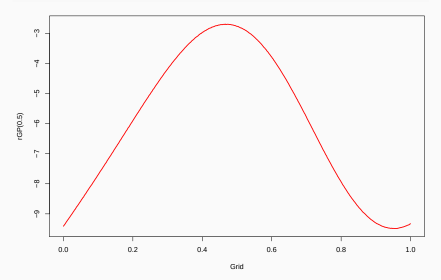




plot(Grid,rGP(0.1),type="l",lwd=2,col="red")



plot(Grid,rGP(0.5),type="1",lwd=2,col="red")

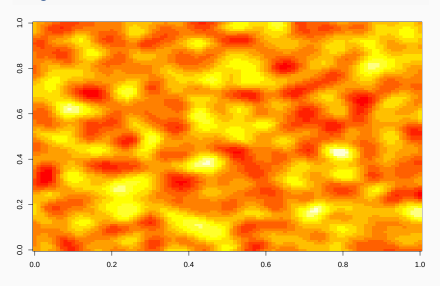


#### 2d GP

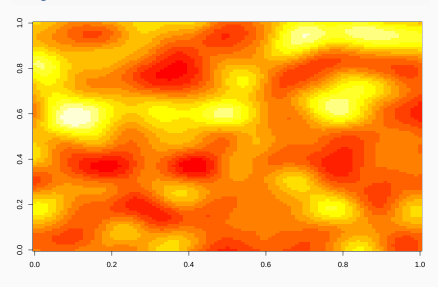
 2d (and 3d) GPs can be useful for modelling smooth spatial heterogeneity

```
Grid = seq(0,1,0.01)
xGrid = expand.grid(Grid,Grid)
Dist = as.matrix(dist(xGrid))
rGP2d = function(r=0.1, s=10) {
   Var = K(Dist,r,s) + (1e-5)*diag(length(Grid)^2)
   z = t(chol(Var)) %*% rnorm(length(Grid)^2)
   matrix(z,nrow=length(Grid))
}
```

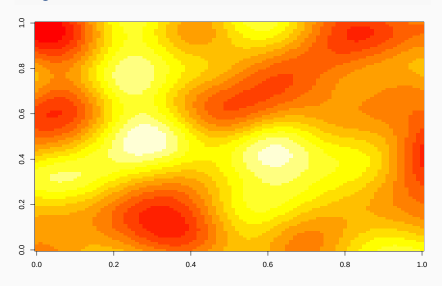
## image(rGP2d(0.05))



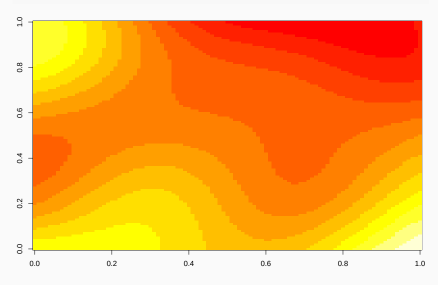
## image(rGP2d(0.1))



## image(rGP2d(0.2))







# Gaussian process emulation

## GP emulation of a computer code

- A GP defines a (tractable) probability distribution over (univariate) functions
- We can condition the GP on some observations and the conditional process is also a GP, but with a new mean and covariance function that is a function of the observations
- This new GP can represent our uncertainty about a function given some observations
- This uncertain representation of our knowledge of the function can be used to *emulate* an expensive function, such as a complex computer code
- The uncertainty associated with the GP can be used to properly quantify and propagage uncertainty associated with the fact that we don't actually know the expensive model everywhere

## **DiceKriging**

 DiceKriging is a popular R package with a range of functionality

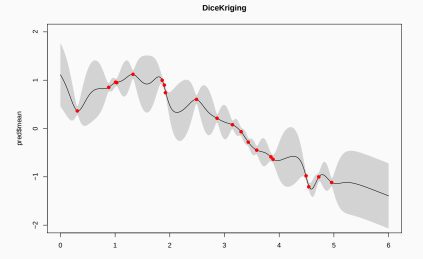
```
library(DiceKriging)
## Loading required package: methods
set.seed(1)
nd = 20
x = as.matrix(runif(nd,0,5))
colnames(x)=c("x")
y = \sin(x) + rnorm(nd, 0, 0.1)
```

```
mod = km(~x,design=x,response=y,nugget=0.001)
##
## optimisation start
## * estimation method : MLE
## * optimisation method : BFGS
## * analytical gradient : used
## * trend model : ~x
## * covariance model :
##
    - type : matern5 2
##
    - nugget : 0.001
##
    - parameters lower bounds : 1e-10
    - parameters upper bounds : 9.301198
##
     - variance bounds : 0.0101135 1.467872
##
##
     - best initial criterion value(s): -10.60975
##
```

```
px = seq(0,6,0.01)
pred = predict(mod,px,type="SK")
```

```
## Warning in checkNames(X1 = X, X2 = newdata, X1.name = ""
## = "newdata"): newdata not named: newdata's variables are
## design
```

```
plot(px,pred$mean,type="l",ylim=c(-2,2),main="DiceKriging")
polygon(c(px,rev(px)),c(pred$upper,rev(pred$lower)),col="l:
lines(px,pred$mean,type="l")
points(x,y,pch=19,col=2)
```



## mlegp

library(mlegp)

## ...done

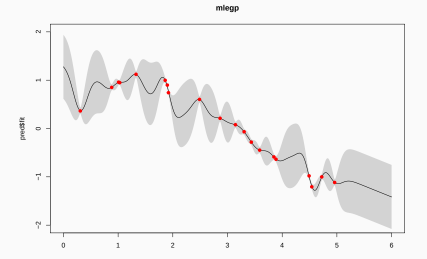
mlegp is an alternative package, based on the book by Santner et al.

```
mod = mlegp(x,y,constantMean=0,nugget.known=1,nugget=0.001)
## no reps detected in meanPerReps, returning originals...
##
## ======= FITTING GP # 1 ================
## running simplex # 1...
## ...done
## ...simplex #1 complete, loglike = 0.895601 (convergence)
## running simplex # 2...
```

## ...simplex #2 complete, loglike = 0.895601 (convergence)

```
pred = predict(mod,as.matrix(px),se.fit=TRUE)
pred$upper = pred$fit + 2*pred$se.fit
pred$lower = pred$fit - 2*pred$se.fit
```

```
plot(px,pred$fit,type="l",ylim=c(-2,2),main="mlegp")
polygon(c(px,rev(px)),c(pred$upper,rev(pred$lower)),col="l:
lines(px,pred$fit,type="l")
points(x,y,pch=19,col=2)
```



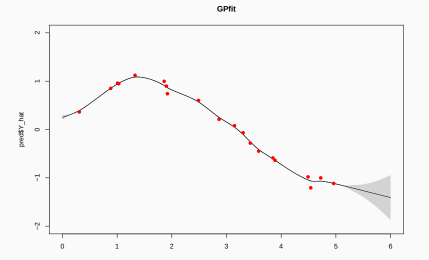
#### **GPfit**

 GPfit is another R package, which has more sophisticated parameter estimation procedures than some other packages

```
library(GPfit)
xScaled = x/5
mod = GP_fit(xScaled,y)
```

```
pxScaled = px/5
pred = predict(mod,as.matrix(pxScaled))
pred$upper = pred$Y_hat + 2*pred$MSE
pred$lower = pred$Y_hat - 2*pred$MSE
```

```
plot(px,pred$Y_hat,type="l",ylim=c(-2,2),main="GPfit")
polygon(c(px,rev(px)),c(pred$upper,rev(pred$lower)),col="l:
lines(px,pred$Y_hat,type="l")
points(x,y,pch=19,col=2)
```



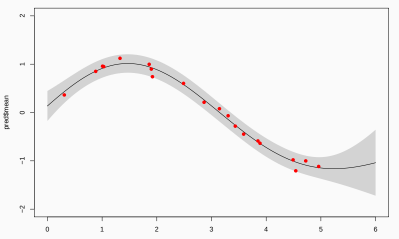
#### hetGP

 hetGP is a modern, sophisticated package for GP emulation, including good support for large datasets, stochastic kriging for noisy models with replication, and heteroskedastic GP modelling

```
library(hetGP)
mod = mleHomGP(x,y) # homoskedastic GP fit
pred = predict(mod,as.matrix(px))
pred$var = pred$sd2 + pred$nugs
pred$sd = sqrt(pred$var)
pred$upper = pred$mean + 2*pred$sd
pred$lower = pred$mean - 2*pred$sd
```

```
plot(px,pred$mean,type="l",ylim=c(-2,2),main="hetGP (mleHor
polygon(c(px,rev(px)),c(pred$upper,rev(pred$lower)),col="l:
lines(px,pred$mean,type="l")
points(x,y,pch=19,col=2)
```



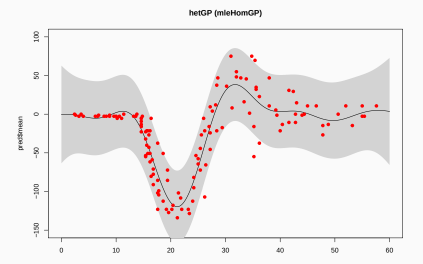


## Heteroskedasticity and the Motorcycle data

- The mcycle dataset from the MASS package is output from a computer model exhibiting heteroskedasticity - non-stationary variance
- We will start with a homoskedastic fit

```
library(MASS)
mod = mleHomGP(mcycle$time,mcycle$accel) # homoskedastic
px=seq(0,60,0.1)
pred = predict(mod,as.matrix(px))
pred$var = pred$sd2 + pred$nugs
pred$sd = sqrt(pred$var)
pred$upper = pred$mean + 2*pred$sd
pred$lower = pred$mean - 2*pred$sd
```

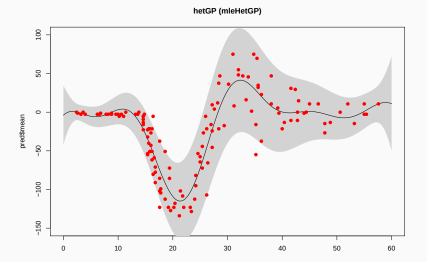
```
plot(px,pred$mean,ylim=c(-150,100),type="l",main="hetGP (main="betGP (main="betGP (main="betGP (main="betGP (main="betGP (pred$upper,rev(pred$lower)),col="laines(px,pred$mean,type="l")
points(mcycle,pch=19,col=2)
```



Next we will do a heteroskedastic fit

```
mod = mleHetGP(mcycle$time,mcycle$accel) # heteroskedastic
pred = predict(mod,as.matrix(px))
pred$var = pred$sd2 + pred$nugs
pred$sd = sqrt(pred$var)
pred$upper = pred$mean + 2*pred$sd
pred$lower = pred$mean - 2*pred$sd
```

```
plot(px,pred$mean,ylim=c(-150,100),type="l",main="hetGP (main="betGP (main="betGP (main="betGP (main="betGP (main="betGP (pred$upper,rev(pred$lower)),col="laines(px,pred$mean,type="l")
points(mcycle,pch=19,col=2)
```



# Design of computer experiments

## Latin hypercube sampling

- When working with expensive computer models with input parameters, the choice of which combination of inputs to run is important, and worth doing carefully
- For a single tunable parameter, a uniform grid can be sensible
- For multiple input parameters, a cartesian product design seems intuitive, but is actually a very poor choice, and gets worse in higher dimensions
- LHS is a technique for choosing points such that the univariate margins essentially form a uniform grid
- The simplest approaches give random designs subject to the constraint of uniform margins
- More sophisticated variants try to optimise some desirable property of the design, such as maximising the minimum distance between points

## The 1hs R package

```
library(lhs)
set.seed(1)

help(package="lhs")
?randomLHS
?maximinLHS
vignette(package="lhs")
vignette("lhs_basics")
```

## A 2d example

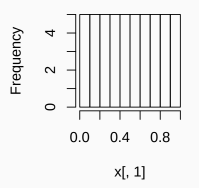
```
x = maximinLHS(50,2)
plot(x, pch=19, col=2)
      0.8
      9.0
x[,2]
      4.0
      0.2
      0.0
                                                               1.0
          0.0
                     0.2
                               0.4
                                          0.6
                                                    8.0
```

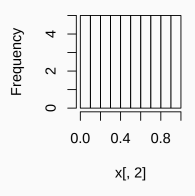
## Univariate margins are uniform

```
op = par(mfrow=c(1,2))
hist(x[,1]); hist(x[,2])
```

## Histogram of x[, 1]

# Histogram of x[, 2]





## Augmenting a design

```
= augmentLHS(x,20)
plot(y,pch=19,col=3); points(x,pch=19,col=2)
     0.8
     9.0
y[,2]
     0.4
     0.2
          0.0
                    0.2
                                        0.6
                                                  8.0
                                                            1.0
                               0.4
```

40

# **Further reading**

#### **Software**

#### R

- CRAN Task Views
  - Spatial
  - Spatio-Temporal
- R Packages
  - hetGP (heteroskedastic GPs)
  - DiceKriging
  - GPfit (robust fitting)
  - mlegp (multiple outputs)
  - Ihs (latin hypercube sampling)

## **Python**

GPy

#### Books and on-line articles

### Wikipedia

- Computer experiment
- Gaussian process
- Latin hypercube sampling
- Uncertainty quantification

#### **Books**

- The Design and Analysis of Computer Experiments (Santner et al.) - ISBN 1493988476
- Gaussian processes for machine learning (Rasmussen and Williams) - ISBN 026218253X