# Creating a sample usage for BACCO from scratch: a cookbook

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## 1 Introduction

Package calibrator of bundle BACCO performs Bayesian calibration of computer models. This document constructs a minimal working example of a simple problem, step by step. Datasets and functions have a .int suffix, representing "intermediate".

This document is not a substitute for KOH or KOHa or Hankin 2005 or the online help files in BACCO. It is not intended to stand alone: for example, the notation used here is that of KOH, and the user is expected to consult the online help in the BACCO package when appropriate.

This document is primarily didactic, although it is informal.

Nevertheless, many of the points raised here are duplicated in the BACCO helpfiles.

Note that many of the objects created in this document are interdependent and changing one sometimes implies changing many others.

The author would be delighted to know of any improvements or suggestions. Email me at r.hankin@noc.soton.ac.uk.

Observations are made over two parameters, x and y, which I style "latitude" and "longitude". I tend to think of the observation as temperature.

The model requires two parameters,  $\hat{A}$  and B and, given these, gives output as a function of x and y. The model is thus a function of four variables: x, y, A, B. Write  $\theta = (A, B)$  to denote the two parameters collectively. KOH use t to denote a specific value of parameters, and  $\theta$  to denote any old set of parameters one might like to consider.

Parameters A and B have true but unknown values and we wish to make inferences about these true values.

# 2 Design matrices

There are two design matrices to consider: D1.int, which is the code run set, and D2.int, the observation points.

In this section, we will generate design matrices of arbitrary size. Data is generated in section 6. First, define the number of code observations (n1) and the number of field observations (n2):

```
> library(calibrator)
> library(emulator)
> n1 <- 20
> n2 <- 21</pre>
```

These can be varied at will. Now create the D1 matrix, of code observation points. This will consist of

```
> D1.int <- latin.hypercube(n1, 4)
> rownames(D1.int) <- paste("coderun", 1:nrow(D1.int), sep = ".")
> colnames(D1.int) <- c("x", "y", "A", "B")
> head(D1.int)
```

```
x y A B coderun.1 0.925 0.025 0.775 0.625 coderun.2 0.825 0.425 0.825 0.225 coderun.3 0.425 0.475 0.525 0.275 coderun.4 0.475 0.225 0.475 0.175 coderun.5 0.575 0.875 0.225 0.925 coderun.6 0.125 0.075 0.625 0.375
```

- Rownames and column-names are applied. They are not strictly necessary, but are very useful in debugging.
- the points are randomly chosen (but fill parameter space reasonably)
- We have ten observation points. More can be added but at the cost of slower run times.

Now the field observation points. This is a two-column matrix, with an  ${\bf x}$  column and a  ${\bf y}$  column.

#### Notes

- See how the number of observations (n1) is different from the number of code runs (n2) as an additional safety check.
- Rownames and columnnames are given.

#### 2.1 Extractor functions

Now we need a function to extract the variable part and the parameter part. Working by analogy to extractor.toy():

#### Notes

- The function returns a two-element list, named x.star and t.vec.
- It just extracts the relevant columns: the first two give the lat and long, and columns three and four give A and B
- drop=FALSE is needed to deal with one-row dataframes consistently.

## 3 Basis functions

We now need basis functions. The basis function h1.toy() is just a constant term with each component. We will use a more sophisticated version.

Recall the fundamental equation of KOH:

$$z(\mathbf{x}) = \rho \eta(\mathbf{x}, \theta) + \delta(\mathbf{x}) + \epsilon \tag{1}$$

where  $z(\mathbf{x})$  is field observation,  $\eta(\cdot, \cdot)$  a Gaussian process with unknown parameters representing a computer model taking two arguments: the first argument is the independent variable  $\mathbf{x}$ , and the second argument a vector of parameters with  $\theta$  being the true but unknown value of the parameters. The  $\delta(\mathbf{x})$  part is a model inadequacy term, also a Gaussian process with unknown parameters. The last term is an observational iid error with  $\epsilon \sim N(0, \lambda^2)$ .

We will take reality to be

$$x + y^2 + y + xy + CE \tag{2}$$

where CE is correlated error. Observations are reality plus uncorrelated  $N(0, \lambda^2)$  error. The model is

$$Ax + By^2 + CE (3)$$

Here the A and B are parameters whose true but unknown values can be determined by comparing equation 2 with equation 3. The values of A and B are thus 1 and 1 respectively; write  $\theta = (A, B)$  to denote the parameters collectively.

The model inadequacy term is thus

$$y + xy + CE$$

Note that this requires A = B = 1. We will take the model basis functions to be

$$h_1(\mathbf{x}, \theta) = (1, x, A, Ax, By^2)^T \tag{4}$$

Notes

• The model output will be

$$h_1(x,\theta)^T \beta_1 + CE$$

where  $\beta_1 = (0, 0, 0, 1, 1)^T$  is the true value of the coefficients; see section 6 for this in use.

• The basis functions could be much more complicated than that or, indeed, much simpler. We could have chosen, for example,  $h_1(x,\theta) = (1, x, y, x^2, y^2, A, B, Ax, Ay, Bx, By, Ax^2, Ay^2, Bx^2, By^2)^T$ ; or, going the other way,  $h_1(x,\theta) = 1$ .

The model inadequacy term then has a mean of y + xy. The basis functions for the model inadequacy is then

$$h_2(x) = (y, xy)^T \tag{5}$$

and the model inadequacy then has a mean of

$$h_2(x)\beta_2$$

where  $\beta_2 = (1,1)^T$ . Again note the great scope for choosing basis functions (although in the case of  $h_2$ () there are no parameters to consider).

### 3.1 Defining regressor functions

```
We need an h1.int():
> h1.int <- function(xin) {
+    out <- c(1, xin[1], xin[3], xin[1] * xin[3], xin[2]^2 * xin[4])
+    names(out) <- c("const", "x", "A", "Ax", "By.sq")
+    return(out)
+ }</pre>
```

Note how the argument xin is just c(x,theta), where x is the independent variables and theta the model parameters. So xin[1] is x, xin[2] is y, xin[3] is A, and xin[4] is B. See how function h1.int() creates the basis functions  $(1, x, A, Ax, By^2)^T$ , as per equation 4.

Now we need a vectorized version:

```
> H1.int <- function(D1) {
+      if (is.vector(D1)) {
+          D1 <- t(D1)
+      }
+      out <- t(apply(D1, 1, h1.int))
+      colnames(out) <- c("const", "x", "A", "Ax", "By.sq")
+      return(out)
+ }</pre>
```

Notes

- Function H1.int is the one needed as arguments to functions like stage1(); function h1.int() is not needed except from within H1.int().
- See how the column names are specified. Again, not strictly necessary but strongly recommended
- See the special consideration for vector D1

Similarly for the model inadequacy function:

```
> h2.int <- function(x) {
+     out <- c(x[1], x[1] * x[2])
+     names(out) <- c("h2.x", "h2.xy")
+     return(out)
+ }
> H2.int <- function(D2) {
+     if (is.vector(D2)) {
+         D2 <- t(D2)
+     }
+     out <- t(apply(D2, 1, h2.int))
+     colnames(out) <- names(h2.int(D2[1, , drop = TRUE]))
+     return(out)
+ }</pre>
```

## 4 Hyperparameter object

We now need a function phi.fun.int() to create an appropriate hyperparameter object. Note that function phi.change() is generic (so we don't need to write one). The best way to do this is to proceed by analogy and modify function phi.fun.toy(), line by line. Most of the function is straightforward to create. Note that the only parts that need changing in this case are the internal functions pdm.maker.psi1() and pdm.maker.psi1(); I use "pdm" to denote "positive

definite matrix". These two functions create positive-definite matrices  $\Omega_x$  and  $\Omega_t$  that are used to determine  $c_1((x,t),(x',t'))$ . These are defined by:

```
> pdm.maker.psi1 <- function(psi1) {
+          jj.omega_x <- diag(psi1[1:2])
+          rownames(jj.omega_x) <- names(psi1[1:2])
+          colnames(jj.omega_x) <- names(psi1[1:2])
+          jj.omega_t <- diag(psi1[3:4], ncol = 2)
+          rownames(jj.omega_t) <- names(psi1[3:4])
+          colnames(jj.omega_t) <- names(psi1[3:4])
+          sigma1squared <- psi1[5]
+          return(list(omega_x = jj.omega_x, omega_t = jj.omega_t, sigma1squared = sigma1squared))
+ }
> pdm.maker.psi2 <- function(psi2) {
+          jj.omegastar_x <- diag(psi2[1:2], ncol = 2)
+          sigma2squared <- psi2[3]
+          return(list(omegastar_x = jj.omegastar_x, sigma2squared = sigma2squared))
+ }</pre>
```

#### Notes

- In function phi.fun.toy() there are two functions with the same name. They extract from vectors psi1 and psi2 a positive definite matrix used in  $c_1(\cdot,\cdot)$  and  $c_2((\cdot,\cdot),(\cdot,\cdot))$ .
- The purpose of all this is to implement the fact that  $\Omega_x$  and  $\Omega_t$  are arbitrary functions of psi1. Here, it's dead easy because the matrices are diagonal and the values just correspond to consecutive elements of psi1 but in general there may be nonzero off-diagonal elements of  $\Omega_x$  which may be a complicated function of psi1.

Given this, we can now change phi.int, using function phi.change(). Note that this function is generic, so we do not need to write another version. The best way to call phi.change() is to define intermediate variables that begin with jj:

```
> jj.psi1 <- 1:5
> names(jj.psi1) <- c("x", "y", "A", "B", "s1sq")
> jj.psi2 <- 1:3
> names(jj.psi2) <- c("x", "y", "s1sq")
> jj.mean1 <- rep(1, 5)
> names(jj.mean1) <- names(jj.psi1)</pre>
> jj.sigma1 <- diag(c(1.1, 1.1, 1.2, 1.3, 1.1))
> rownames(jj.sigma1) <- names(jj.psi1)</pre>
> colnames(jj.sigma1) <- names(jj.psi1)</pre>
> jj.mean2 <- c(1, 0.1, rep(1.1, 3))
> names(jj.mean2) <- c("rho", "lambda", names(jj.psi2))
> jj.sigma2 <- diag(c(1, 0.2, 1.1, 1.1, 1.2))/10</pre>
> rownames(jj.sigma2) <- names(jj.mean2)</pre>
> colnames(jj.sigma2) <- names(jj.mean2)</pre>
> jj.mean.th <- 1:2
> names(jj.mean.th) <- c("A", "B")
> jj.sigma.th <- diag(c(1.5, 1.7))
> rownames(jj.sigma.th) <- names(jj.mean.th)</pre>
> colnames(jj.sigma.th) <- names(jj.mean.th)</pre>
```

These variables may be passed directly to phi.fun.int():

```
> phi.int <- phi.fun.int(rho = 1, lambda = 1, psi1 = jj.psi1, psi2 = jj.psi2,
+ psi1.apriori = list(mean = jj.mean1, sigma = jj.sigma1),
+ psi2.apriori = list(mean = jj.mean2, sigma = jj.sigma2),
+ theta.apriori = list(mean = jj.mean.th, sigma = jj.sigma.th),
+ power = 2)</pre>
```

- The purpose of the call to phi.fun.int() was to create a hyperparameter object, here phi.int.
- you only have to call this function, with all its complicated arguments, once. Save the result in, say, jj and then use function phi.change(), which is generic, and much simpler to use.
- The value for rho and lambda and power are just scalars
- psi1 and psi2 are structures, or named vectors (it's always good practice to elements: two for x and y, two for the parameters A and B, and one for sigma1squared. Vector psi2 has three: two for x and y, and one for sigma2squared.
- the priors are as expected. Remember that the apriori distributions are over psi1 or psi2, not just the roughness lengths.

So now we have a working hyperparameter object phi.int, we can modify it with generic function phi.change():

```
> phi.int2 <- phi.change(old.phi = phi.int, phi.fun = phi.fun.int,
+ rho = 3)
> print(phi.int2$rho)
[1] 3
```

Note that function phi.change() takes an old.phi argument, which is a working hyperparameter object to be modified. It also takes a phi.fun argument, which is the name of a hyperparameter creation function, in this case phi.fun.int.

# 5 Functions E.theta.int() and Edash.theta.int()

The online help page for E.theta.toy() discusses how to create a new function for use in a particular example.

We need to define functions E.theta.int() and Edash.theta.int(). These functions change when the basis functions h1.int() and h2.int() change, because of possible nonlinearity.

The function E.theta.toy() is relatively simple because there the basis functions were linear in  $\theta$ , so in this case expectation commutes past taking the basis functions<sup>1</sup>

The first step is to examine function E.theta.toy():

```
> E.theta.toy

function (D2 = NULL, H1 = NULL, x1 = NULL, x2 = NULL, phi, give.mean = TRUE)
{
   if (give.mean) {
      m_theta <- phi$theta.apriori$mean
      return(H1(D1.fun(D2, t.vec = m_theta)))
   }</pre>
```

<sup>&</sup>lt;sup>1</sup>This is not true in general. Consider a nonlinear case, for example  $h_1(x,\theta) = (1,A^2x)^T$ . Now  $E_{\theta}(h_1(x,\theta))^T$ —that is, the expectation of  $(1,A^2x)^T$  under the prior distribution for  $\theta$ —will be  $\left(1,(\overline{A}^2+\sigma_A^2)x\right)^T$ .

```
else {
    out <- matrix(0, 6, 6)
    out[4:6, 4:6] <- phi$theta.apriori$sigma
    return(out)
  }
}
</pre>

    The difficulty is that the basis functions are nonlinear Re
```

The difficulty is that the basis functions are nonlinear. Recall that function h1.int() includes a product:

```
> h1.int
function (xin)
{
   out <- c(1, xin[1], xin[3], xin[1] * xin[3], xin[2]^2 * xin[4])
   names(out) <- c("const", "x", "A", "Ax", "By.sq")
   return(out)
}</pre>
```

We can use the fact that  $E(XY) = E(X) \cdot E(Y)$  if X and Y are independent random variables. As the prior variance matrix (viz, jj.psil.apriori) is diagonal, different terms are independent. BUT,  $E(X^2) \neq (E(X))^2$ . Thus we may copy from function E.theta.toy() and modify as necesary:

```
> E.theta.int <- function(D2 = NULL, H1 = NULL, x1 = NULL, x2 = NULL,
      phi, give.mean = TRUE) {
      if (give.mean) {
+
          m_theta <- phi$theta.apriori$mean</pre>
          return(H1(D1.fun(D2, t.vec = m_theta)))
      }
      else {
          out <- matrix(0, 5, 5)
          out[3, 3] <- phi$theta.apriori$sigma[1, 1]</pre>
          out[3, 4] \leftarrow phi$theta.apriori$sigma[1, 1] * x1[1]
          out[4, 3] <- phi$theta.apriori$sigma[1, 1] * x2[1]
          out[4, 4] <- phi$theta.apriori$sigma[1, 1] * x1[1] *
               x2[1]
          out[5, 5] <- phi$theta.apriori$sigma[2, 2] * x1[2] *
              x2[2]
          return(out)
      }
+ }
```

Notes

- The object returned when give.mean is TRUE does not need changing, because h1.int() is linear in A and B. If there were nonlinear terms there (such as  $A^2$ ) we would need to add a variance.
- The online help for E.theta.toy() discusses the give.mean being FALSE part. Here, we use the fact that the prior distribution has zero correlation between A and B. The thing at the top of page 5 of the supplement should be

$$E_{\theta}\left(\mathbf{h}_{1}\left(\mathbf{x}_{j},\boldsymbol{\theta}\right)\mathbf{h}_{1}\left(\mathbf{x}_{j},\boldsymbol{\theta}\right)^{T}\right) = E_{\theta}\begin{pmatrix} 1 & x & A & Ax & By^{2} \\ x & x^{2} & Ax & Ax^{2} & Bxy^{2} \\ A & Ax & A^{2} & A^{2}x & ABy^{2} \\ Ax & Ax^{2} & A^{2}x & A^{2}x^{2} & ABxy^{2} \\ By^{2} & Bxy^{2} & ABy^{2} & ABxy^{2} & B^{2}y^{4} \end{pmatrix}$$
(6)

which has value

$$\begin{pmatrix} 1 & x & \overline{A} & \overline{A}x & \overline{B}y^2 \\ x & x^2 & \overline{A}x & \overline{A}x^2 & \overline{B}xy^2 \\ \overline{A} & \overline{A}x & \overline{A}^2 + \sigma_A^2 & \left(\overline{A}^2 + \sigma_A^2\right)x & \left(\overline{A}\overline{B} + \cos(A, B)\right)y^2 \\ \overline{A}x & \overline{A}x^2 & \left(\overline{A}^2 + \sigma_A^2\right)x & \left(\overline{A}^2 + \sigma_A^2\right)x^2 & \left(\overline{A}\overline{B} + \cos(A, B)\right)xy \\ \overline{B}y^2 & \overline{B}xy^2 & \left(\overline{A}\overline{B} + \cos(A, B)\right)y^2 & \left(\overline{A}\overline{B} + \cos(A, B)\right)xy^2 & \left(\overline{B}^2 + \sigma_B^2\right)y^4 \end{pmatrix}$$

and so the object to return (when give.mean is FALSE) is this, minus  $E_{\theta}(h_1(x,\theta)) E_{\theta}(h_1(x,\theta))^T$  as documented. This would be

$$\begin{pmatrix}
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & \sigma_A^2 & \sigma_A^2 x & \cos(A, B)y^2 \\
0 & 0 & \sigma_A^2 x & \sigma_A^2 x^2 & \cos(A, B)xy \\
0 & 0 & \cos(A, B)y^2 & \cos(A, B)xy & \sigma_B^2 y^4
\end{pmatrix}$$

And remembering that the covariance is zero this reduces to

$$\begin{pmatrix}
0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 \\
0 & 0 & \sigma_A^2 & \sigma_A^2 x & 0 \\
0 & 0 & \sigma_A^2 x & \sigma_A^2 x^2 & 0 \\
0 & 0 & 0 & 0 & \sigma_R^2 y^4
\end{pmatrix}$$

which is implemented in function E.theta.int() given above<sup>2</sup>

• There is probably a better way to return this matrix.

#### 5.1 Function Edash.theta.int()

Function Edash.theta.int() returns expectation of  $h_1(x,\theta)$  with respect to the normal distribution

$$\mathcal{N}\left( (\boldsymbol{V}_{\theta}^{-1} + 2\boldsymbol{\Omega}_{t})^{-1} (\boldsymbol{V}_{\theta}^{-1} \boldsymbol{\Omega}_{\theta} + 2\boldsymbol{\Omega}_{t} \mathbf{t}_{k}), (\boldsymbol{V}_{\theta}^{-1} + 2\boldsymbol{\Omega}_{t})^{-1} \right)$$
(7)

Fortunately,  $E_{\theta}(h \cdot h^T)$  is not required.

As the basis functions used are linear in theta<sup>3</sup>, expectation WRT the dashed distribution commutes past the regressor function  $h_1(\cdot,\cdot)$ . So we can just use the toy function:

#### > Edash.theta.int <- Edash.theta.toy

<sup>2</sup>Returning to the nonlinear example given in the previous footnote, viz  $h = h_1(x, \theta) = (1, A^2x)^T$ . This would

have 
$$E_{\theta}(h) = \left(1, x(\overline{A}^2 + \sigma_A^2)\right)^T$$
 and  $E_{\theta}(h) \cdot E_{\theta}(h)^T = \begin{pmatrix} 1 & x\left(\overline{A}^2 + \sigma_A^2\right) \\ x\left(\overline{A}^2 + \sigma_A^2\right) & x^2\left(\overline{A}^2 + \sigma_A^2\right) \end{pmatrix}$ .

But what is needed by KOH is

$$E_{\theta}\left(h \cdot h^{T}\right) = E_{\theta}\left(\begin{array}{cc} 1 & A^{2}x \\ A^{2}x & A^{4}x^{2} \end{array}\right) = \left(\begin{array}{cc} 1 & x\left(\overline{A}^{2} + \sigma_{A}^{2}\right) \\ x\left(\overline{A}^{2} + \sigma_{A}^{2}\right) & x^{2}\left(\overline{A}^{4} + 6\overline{A}^{2}\sigma_{A}^{2} + 3(\sigma_{A}^{2})^{2}\right) \end{array}\right).$$

The manpage for function E.theta.toy() says how, if argument give.mean is FALSE, the value returned should be the thing that has to be added to  $E_{\theta}(h) \cdot E_{\theta}(h)^T$  to give  $E_{\theta}(h \cdot h^T)$ . The motivation for considering it this way is that E.theta.toy(...,give=FALSE) will return a zero matrix for basis functions linear in  $\theta$ .

So, for this  $h(\cdot,\cdot)$ , function E.theta.int() would have to return the difference between  $E_{\theta}\left(h\cdot h^{T}\right)$  and  $E_{\theta}(h)\cdot$ 

$$E_{\theta}(h)^T$$
, which would be  $\begin{pmatrix} 0 & 0 \\ 0 & x^2 \left( 4\overline{A}^2 \sigma_A^2 + 2(\sigma_A^2)^2 \right) \end{pmatrix}$ .

<sup>3</sup>Returning to our nonlinear example, footnotes passim, viz  $h_1(x,\theta) = (1,xA^2)^T$ , we can see from equation 7

## 6 Data

We now generate some data: code runs, and observations.

This is done in order to test the routines: by generating data with known parameters and hyperparameters we can verify that the package can reproduce at least approximately correct values.

Consider the following equation, taken from KOH:

$$z_i = \zeta(x_i) + e_i = \rho \eta(x_i, \theta) + \delta(x_i) + e_i \tag{8}$$

where  $z_i$  is the *i*-th observation,  $\zeta(x_i)$  is the true value at point  $x_i$ ,  $\rho$  a calibration factor (of notional value 1),  $\eta(\cdot, \cdot)$  the code viewed as a function of observation point and parameter value,  $\theta$  the true but unknown set of parameters, and  $\delta(\cdot)$  the model inadequacy term, and  $e_i \sim \mathcal{N}(0, \lambda^2)$  is an observational error term.

So, do the model first. Recall that the model is a Gaussian process and we have discussed the mean in section 3. The variance matrix is given by function corr.matrix() of package emulator. We know,  $ex\ cathedra$ , that A=B=1:

```
> theta.TRUE <- c(1, 1)
```

and we can specify beta1 and psi1:

```
> beta1.TRUE <- c(0, 0, 0, 1, 1)
> psi1.TRUE <- c(4, 4, 4, 4, 0.5)
```

Now we need to create a design matrix:

```
> two.designs <- rbind(D1.int, D1.fun(x.star = D2.int, t.vec = theta.TRUE))
```

Here, two.designs is a design matrix of n1+n2 rows; the first n1 rows of which are the code observation points, and the last n2 rows are the field observation points but with the true parameter value added.

Just have a look at the first three and last three lines:

```
> two.designs[c(1:3, (n1 + n2):(n1 + n2 - 2)),]
```

```
x y A B coderun.1 0.9250000 0.02500000 0.775 0.625 coderun.2 0.8250000 0.42500000 0.825 0.225 coderun.3 0.4250000 0.47500000 0.525 0.275 obs.21 0.8809524 0.07142857 1.000 1.000 obs.20 0.4523810 0.92857143 1.000 1.000 obs.19 0.7857143 0.30952381 1.000 1.000
```

See how the block of 1.0s in the lower right corner corresponds to appending the true value of  $\theta$  to the design matrix.

The next step is to sample from the appropriate multivariate Gaussian distribution:

```
> jj.mean <- H1.int(two.designs) %*% beta1.TRUE
> jj.sigma <- psi1.TRUE[5] * corr.matrix(two.designs, scales = psi1.TRUE[1:4])
> code.and.obs <- as.vector(rmvnorm(n = 1, mean = jj.mean, sigma = jj.sigma))
> y.int <- code.and.obs[1:n1]
> z.int <- code.and.obs[(n1 + 1):(n1 + n2)]
> names(y.int) <- rownames(D1.int)
> head(y.int)
```

that  $E'_{\theta} = \left(1, x\left({\mu'}_A^2 + {\sigma'}_A^2\right)\right)^T$  where  $\mu'_A$  is the mean of the distribution in equation 7, that is, the first element of the vector  $(\boldsymbol{V}_{\theta}^{-1} + 2\Omega_t)^{-1}(\boldsymbol{V}_{\theta}^{-1}\Omega_{\theta} + 2\Omega_t t_k)$ , and  ${\sigma'}_A^2$  is the variance—that is the top left element of  $(\boldsymbol{V}_{\theta}^{-1} + 2\Omega_t^{-1})$  (NB: this argument is true whether or not the variance matrix  $\boldsymbol{V}_{\theta}$  and the positive definite scales matrix  $\Omega_t$  are diagonal)

```
coderun.1 coderun.2 coderun.3 coderun.4 coderun.5 coderun.6 0.8361985 -0.7668183 -0.9380283 -1.5809861 -0.6440748 -1.1132955
```

- The mean, jj.mean, is given by the linear combination of the regressor basis as discussed in section 3
- The model  $\eta$  is a Gaussian process. The mean is specified, the variance matrix given by the correlation function corr.matrix() of package emulator. The value for  $\sigma_1^2$  is one (ie the fifth element of psi1.TRUE).
- Function rmvnorm() returns a matrix, which has to be converted to a vector.
- The names of y.int have to be specified explicitly.
- Observe how n=1 in the call to rmvnorm(). We are making a *single* observation of a multivariate Gaussian distribution.

## 6.1 Observations and model inadequacy

To create reality, we have to generate model observations using the true but unknown parameter values.

We now have to add the model inadequacy term to z.int. First, specify the parameters and hyperparameters of the model inadequacy:

```
> beta2.TRUE <- c(1, 1)
> psi2.TRUE <- c(3, 3, 0.6)
```

Which give the true values. Now create model inadequacy:

```
> jj.mean <- drop(H2.int(D2.int) %*% beta2.TRUE)
> jj.sigma <- corr.matrix(D2.int, scales = psi2.TRUE[1:2]) * psi2.TRUE[3]
> model.inadequacy <- rmvnorm(n = 1, mean = jj.mean, sigma = jj.sigma)
> z.int <- as.vector(z.int + model.inadequacy)
> names(z.int) <- rownames(D2.int)</pre>
```

Notes

- The overall purpose of the above code fragment is to create observations drawn from the appropriate multivariate Gaussian distribution. The key is the fourth line, in which the model observations cond.gp have model inadequacy added.
- Model inadequacy has two components: the mean (jj.mean), generated using the true coefficients and the basis functions; and the correlated residual.
- The correlated error is added using rmvnorm() with a variance matrix generated by corr.matrix().
- The correlated error is independent of the model runs, and in particular is not a function of  $\theta$ .

Now add observational error:

```
> lambda.TRUE <- 0
> jj.obs.error <- rnorm(n2) * lambda.TRUE
> z.int <- z.int + jj.obs.error
> head(z.int)

    obs.1    obs.2    obs.3    obs.4    obs.5    obs.6
-0.2952739    1.0991959 -0.2293498 -1.9314573 -0.2258510    1.5699037
```

- The observation errors are uncorrelated Gaussian with a mean of 0 and a variance of 0.1.
- Thus  $\lambda$  is 0.1

And finally we need to create a full data vector d.int

```
> d.int <- c(y.int, z.int)
```

## 7 Intermediate results

We now use some functions that are part of the BACCO bundle.

Note that the numbers given above will vary from instantiation to instantiation.

OK, so let's change the hyperparameter object to contain the true values:

```
> phi.true <- phi.change(phi.fun = phi.fun.int, old.phi = phi.int,
+ psi1 = psi1.TRUE, psi2 = psi2.TRUE, lambda = 0.1, rho = 1)
and then use these hyperparameters to estimate the coefficients:</pre>
```

```
> betahat.fun.koh(theta = theta.TRUE, d = d.int, D1 = D1.int, D2 = D2.int,
+ H1 = H1.int, H2 = H2.int, phi = phi.true)
```

```
[,1]
const -0.6869341
x -0.1125843
A -0.4563367
Ax 2.1940675
By.sq 0.8846170
h2.x 1.5769657
h2.xy -0.6339745
```

(note the last argument to betahat.fun.koh()).

We know the correct answer should be c(beta1.TRUE,beta2.TRUE)=c(0,0,0,1,1,1,1,1), so there is evidently a lot of scatter. Try redefining psi1.TRUE and psi2.TRUE so that the variance (ie the last element of each vector) is smaller. Or for that matter, using a larger value of n1 in section 2.

Just as a point of interest, we will estimate the betas but using the wrong parameters (recall that in practice, the parameters' true value is not known).

This is far from the true values because we have used a very inaccurate value for theta.

#### 7.1 Calibration

As a bit of fun, we can use equation 8 of the supplement, implemented in BACCO by p.eqn8.supp(). This equation gives the probability of theta, given d and psi. Note that this formula is conditional on the correct hyperparameters psi, so we will use phi.true.

```
> p.eqn8.supp(theta = c(1, 1), D1 = D1.int, D2 = D2.int, H1 = H1.int, + H2 = H2.int, d = d.int, phi = phi.true)
```

[1] 1849.819

(recall that this function gives a number *proportional* to the true probability). Now try a different value for theta:

```
> p.eqn8.supp(theta = c(5, -6), D1 = D1.int, D2 = D2.int, H1 = H1.int,
+ H2 = H2.int, d = d.int, phi = phi.true)
```

See how this is lower, because the value of theta is unlikely to be the true one. So we could use a maximum likelihood estimator (which would have to use numerical optimization techniques) to estimate theta, if we knew the correct values to use for the hyperparameters.

In practice, the hyperparameters are not known in advance. Estimating them is the subject of the next section.

## 8 Estimating the hyperparameters

The BACCO bundle includes two functions to automate the determination of the hyperparameters: stage1() and stage2(). The bundle also includes a stage3() function which gives a MLE for theta (but no such stage3 appears in KOH).

The function calls in this section are very computationally intensive and the calls include a number of timesaving features (such as very short optimization) that degrade the quality of the predictions.

These features are discussed where appropriate but the user is advised to fiddle with them to get better results. YMMV.

#### 8.1 Stage 1

KOH proposed estimating the hyperparameters in two stages. Stage 1 used just the code output data y to estimate psi1. In the calibrator bundle, this is accomplished by stage1():

```
> phi.stage1 <- stage1(D1 = D1.int, y = y.int, H1 = H1.int, maxit = 10,
+ method = "SANN", trace = 0, do.print = FALSE, phi.fun = phi.fun.int,
+ phi = phi.int)</pre>
```

Notes

- Function stage1() returns a hyperparameter object, which contains optimized value for psi1.
- The method used is simulated annealing, because with maxit=1 it finishes very quickly.
- The function takes a hyperparameter object, in this case phi.int.
- By default, function stage1() maximizes the posterior probability, so the prior (which is part of the hyperparameter object) makes a difference.

We can examine the output of stage1() directly:

```
> phi.stage1$psi1
                                                         s1sq
3.5507174 0.8593747 1.1535640 5.8098033 3.3226697
   Recall that phi.stage1$psi1 is a vector of free parameters that are used to calculate c_1(\mathbf{x}, \mathbf{x}').
   Further recall that c_1((\mathbf{x}, \mathbf{t}), (\mathbf{x}', \mathbf{t}')) = \sigma_1^2 \exp\left\{-(\mathbf{x} - \mathbf{x}')^T \mathbf{\Omega}_x(\mathbf{x} - \mathbf{x}') - (\mathbf{t} - \mathbf{t}')^T \mathbf{\Omega}_t(\mathbf{t} - \mathbf{t}')\right\}
where \sigma_1^2, and \Omega_x and \Omega_t are positive definite matrices that are functions of psi1.
   In this case we have
> phi.stage1$sigma1squared
     s1sq
3.322670
> phi.stage1$omega_x
x 3.550717 0.0000000
y 0.000000 0.8593747
> phi.stage1$omega_t
A 1.153564 0.000000
B 0.000000 5.809803
8.2
       Stage 2
Stage 2 is the estimation of \rho, \lambda, and psi2. This is accomplished by function stage().
   NB: stage 2 is very computationally intensive!
   OK, use stage2():
> use1 <- 1:10
> use2 <- 1:11
> phi.stage2 <- stage2(D1 = D1.int[use1, ], D2 = D2.int[use2, ],
       H1 = H1.int, H2 = H2.int, y = y.int[use1], z = z.int[use2],
       extractor = extractor.int, phi.fun = phi.fun.int, E.theta = E.theta.int,
       Edash.theta = Edash.theta.int, maxit = 1, method = "SANN",
       phi = phi.stage1)
[1] 0.0000000 0.0000000 0.0000000 0.6931472 1.0986123
[1] -42.93561
sann objective function values
initial
                 value 42.935605
```

#### Notes

final

value 42.935605

sann stopped after 0 iterations

- This function takes a long long long time to run, and is very computationally slow.
- The function uses only the first 10 code runs and the first 11 field observations. You can change this by modifying use1 and use2 in the above chunk.

- The above function call is an absolute minimum working model. It uses SANN with only one function evaluation.
- The start point is the output from stage1(), viz phi.stage1.
- The purpose of function stage2() is to optimize the posterior probability of the hyperparameters rho, lambda, sigma2squared, and the lengthscales for  $c_2(\cdot,\cdot)$ .

The modified hyperparameters are

```
> phi.stage2$rho
1
> phi.stage2$lambda
1
> phi.stage2$sigma2squared
s1sq
    3
> phi.stage2$psi2
    x    y s1sq
    1    2    3
```

(although, given the extremely short optimization run above, these parameters may well not be different from the original).

# 9 Calibrated prediction

Function EK.eqn10.supp() carries out calibrated prediction as per section 4.2 of KOH2. It should come as no surprise that

- The R code is complicated and impenetrable
- The function arguments are tedious, error-prone, and complicated
- The mathematics are complicated and impenetrable
- The numerics take an interminably long time to complete
- Even with the simplest case possible, accurate results require more computing power than is available in the entire universe.

```
... but, it is implemented.
```

The first step is to read help(EK.eqn10.supp), which gives a working example.

There are two arguments that are not covered above. The first is X.dist and the second is hbar. These are discussed in the online help pages.

For X.dist, we need to define an uncertainty distribution on the independent variables. Working from X.dist.toy, we may copy it directly:

```
> jj.xdist.mean <- rep(0.5, 2)
> names(jj.xdist.mean) <- c("x", "y")
> jj.xdist.var <- 0.05 + diag(c(0.1, 0.1))
> rownames(jj.xdist.var) <- c("x", "y")
> colnames(jj.xdist.var) <- c("x", "y")
> X.dist.int <- list(mean = jj.xdist.mean, var = jj.xdist.var)</pre>
```

thus defining a Gaussian uncertainty distribution for X. We now need a function hbar. The manpage discusses this and gives an example. Note that the example is excruciatingly simple because there the basis functions are linear in x. And in the intermediate case they are not.

To write a suitable function, we need to remind ourselves what the basis functions h1.int() and h2.int() are. Look back at equations 4 and 5.

Take the top bit first. This is  $\rho E_X \{h_1(\mathbf{x}, \boldsymbol{\theta})\}$ , that is, expectation with respect to X.dist. Recall that  $h_1(\mathbf{x}, \boldsymbol{\theta}) = (1, x, A, Ax, By^2)^T$ , so the top bit is

$$\rho E_X \left( 1, x, A, Ax, By^2 \right)^T = \rho \left( 1, \overline{x}, A, A\overline{x}, B \left( \overline{y}^2 + \sigma_y^2 \right) \right)^T \tag{9}$$

where the overline refers to expectation with respect to X.dist. The bottom bit is  $E_X \{h_2(\mathbf{x})\}$ , which is (recall that  $h_2(\mathbf{x}) = (x, xy)^T$ ):

$$E_X(x, xy)^T = (\overline{x}, \overline{x} \cdot \overline{y} + \text{cov}(x, y))^T$$
(10)

and we can use the fact that  ${\tt X.dist.int}$  specifies zero correlation between the two independent variables' uncertainty distribution to reduce this to

$$(\overline{x}, \overline{x} \cdot \overline{y})^T$$

OK, using the toy example as a guide, we can define a working hbar.fun.int():

```
> hbar.fun.int <- function(theta, X.dist, phi) {
+      if (is.vector(theta)) {
+          theta <- t(theta)
+      }
+      first.bit <- phi$rho * H1.int(D1.fun(X.dist$mean, theta))
+      first.bit[, 5] <- first.bit[, 5] + theta[, 2] * X.dist$var[2,
+          2]
+      second.bit <- H2.int(X.dist$mean)
+      jj.names <- colnames(second.bit)
+      second.bit <- kronecker(second.bit, rep(1, nrow(first.bit)))
+      colnames(second.bit) <- jj.names
+      return(t(cbind(first.bit, second.bit)))
+ }</pre>
```

notes

- See how the fifth column of first.bit has to have a component for the variance added, to match the variance term in equation 9.
- There is no such addition to **second.bit** because the random variables in **X.dist.int** have zero correlation.
- Everything else just follows from hbar.fun.toy().

Now we can conduct a calibrated prediction:

```
> jj <- EK.eqn10.supp(X.dist = X.dist.int, D1 = D1.int, D2 = D2.int,
+ H1 = H1.int, H2 = H2.int, d = d.int, hbar.fun = hbar.fun.int,
+ lower.theta = c(-3, -3), upper.theta = c(3, 3), extractor = extractor.int,
+ phi = phi.stage2, maxpts = 10)
[1] TRUE
> jj
[1] 0.3300762
```

- $\bullet$  We use the optimized value for phi, that is phi.stage2.
- I have set maxpts to 10, as adapt() is very slow with the default settings. We could also fiddle with the limits of integration, viz theta.lower and theta.upper.

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