## Model Calibration with the FME package

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### Foreword

This is an R Markdown document. Markdown is a simple formatting syntax for authoring PDF documents.

When you click the **Knit** button in Rstudio a pdf will be generated that includes both content as well as the output of any embedded R code chunks within the document. You can then modify the code inside the r chunk and update the pdf accordingly.

#### **Objectives**

This is an example of script as should be produced during practical session of the lecture "Modélisation des écosystèmes et des cycles biogéochimiques" (Partim: Resource competition). It consist in calibrating the parameter of a simple growth function, in order to render observation of growth rates measured at different resource availability. (Those data have been created artificially and are located in a text file). To do so we will exploit functions from "FME" library (Soetaert and Petzoldt 2010). The notions we are going to consider are pretty well explained in this documentation. To start, we simply load the FME library (which will automatically load the dependence libraries).

#### library(FME)

```
## Loading required package: deSolve
## Loading required package: rootSolve
## Loading required package: coda
```

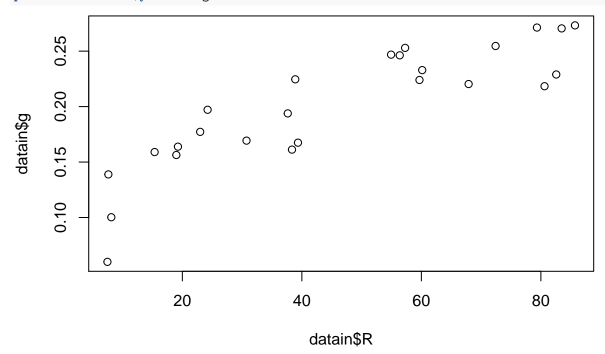
## Load Data

Now we load the data using the read.table function which store them in a "data frame" structure (datain). We then assign names to the columns of datain. Remember the command c("R", "g") create a vector of two string elements: "R" and "g". Here R goes for ressource availability and g for corresponding growth rates. To assign column names, you can use the function colnames

```
datain<-read.table(file="G1R1.txt")
colnames(datain)<-c("R", "g")</pre>
```

Let us have a first view on those data and plot them (use plot(x=, y=))

plot( x=datain\$R ,y=datain\$g)



## Build a growth model

The next block defines the function Growth using a simple Michalis-Menten function form

$$g = g_{max} \frac{R}{R+k}$$

Growth takes two arguments:

- pars is a vector of two parameter:  $gM(g_{max})$  is the maximum growth rate, and k is the constant which determines how strongly the resource availability restricts growth (when R = k,  $g = g_{max}/2$ ).
- R is the resource availability

The function returns a data frame with the ressource values R, and the corresponding growth rates g.

```
Growth <-function (pars,R) {
  with(as.list(pars),{
    g <- gmax*(R / (R + k ))
    return (data.frame(R=R,g=g))
  })
}</pre>
```

Now we define a vector of parameter parms with guess values (try k=10 and gmax= .8) and will use it to plot a curve depicting how the growth rate vary with the resource availability. Note that the growth rates are obtained using the function Growth previously defined, and depend on the set of parameter we are passing when calling this function.

The function Growth returns a data frame with both input R and resulting g. This is why in the following we have to use the nomenclature growthoutput\$g the extract the column g from the data frame growthoutput.

To Do

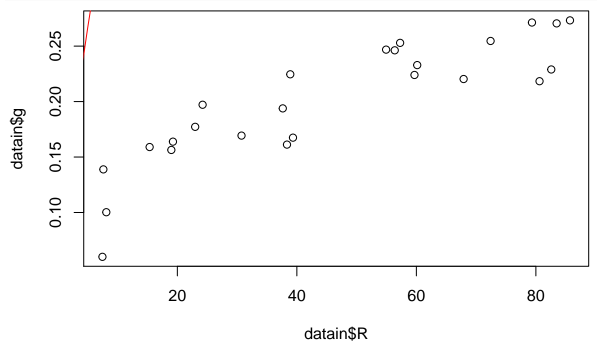
- Define the parameter vector
- Call the growth function, using this parameters, for a range of R=seq(0,100,.5)
- plot the points (as previoulsy) and add the model prediction, using the command lines (use col=red as argument)

```
parms <- c(k = 10, gmax=.8)

# Plot data points from the text file
plot( x=datain$R ,y=datain$g)

growthoutput=Growth(parms,seq(0,100,.5))

# Plot computed growth rates for a range R going from 0 to 100 by step of .5
lines(x=growthoutput$R,y=growthoutput$g, col="red")</pre>
```



## Calibration

To adjust the parameters we first to quantify the model misfit, ie. we need a function that receives a set of parameter and returns the residuals (ie, the difference between the g measured and those predicted by the growth model at the R value considered in the measurements).

```
ModelCosta <- function(p) {
  out <- Growth(p, datain$R)</pre>
```

```
return( datain$g-out$g ) # residuals
}
ModelCosta(parms)

## [1] -0.4612047 -0.2580135 -0.4281564 -0.4391865 -0.4332666 -0.4933549

## [7] -0.2070433 -0.4482894 -0.4343023 -0.4769833 -0.4704252 -0.4530495

## [13] -0.3691216 -0.4733228 -0.3255556 -0.4431999 -0.3802238 -0.4439216

## [19] -0.4846442 -0.3627637 -0.3676948 -0.4118085 -0.4299545 -0.4382192

## [25] -0.2818159
```

We can now use the function modFit, to find the set of parameters that minimize the residuals given by ModelCosta. Here, we selected the "Pseudo" search algorithm and imposed bounds for the parameters [0 1000] and [0 1000] (type '?modFit' for more Details).

Fita is now an object of the class modFit, which contains a number of attributes, and which can be used as an argument to dedicated functions as illustrated below

- summary gives some stats on the calibration procedure
- coef returns the value of the best parameters

```
\# What is the value of the fitted parameters, and the standard error on those parameters ? \operatorname{summary}(\operatorname{Fita})
```

```
##
## Parameters:
        Estimate Std. Error t value Pr(>|t|)
##
## k
        16.68208
                    3.39933
                              4.907 5.87e-05 ***
## gmax 0.29843
                    0.01736 17.191 1.27e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.02401 on 23 degrees of freedom
##
## Parameter correlation:
##
             k
                 gmax
        1.0000 0.9163
## gmax 0.9163 1.0000
coef(Fita)
##
            k
                    gmax
```

We can now check visually that the fitted parameters provides a better description of the data, by evaluating  ${\tt Growth}$  on the same range of R but using the new parameters.

To Do

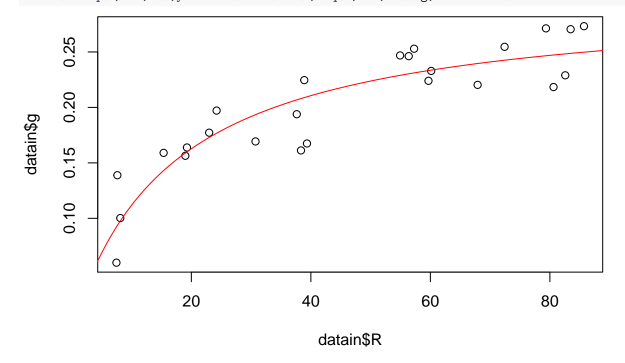
## 16.6820785

• Start by plotting the data, as previously.

0.2984294

Now use lines command to show model outputs, but provides the best parameters from the modFit
procedure to evaluate the model

# # Plot plot(x=datain\$R,y=datain\$g) lines(x=seq(0,100,.5),y=Growth(coef(Fita),seq(0,100,.5))\$g, col="red")



## Sensitivity

Which parameter has the more impact on the model output ?

The function sensFun evaluates sensitivity function around the prescribed parameter values (type ?sensFun for definitions).

$$S_{i,j} = \frac{\partial y_i}{\partial \Theta_j}$$

where,

- $y_i$  is the output variable (g in our case)
- $\Theta_j$  is a parameter

These value can be summarized using the scales :

$$L_1(\Theta_j) = \sum |S_{i,j}|/n$$

and

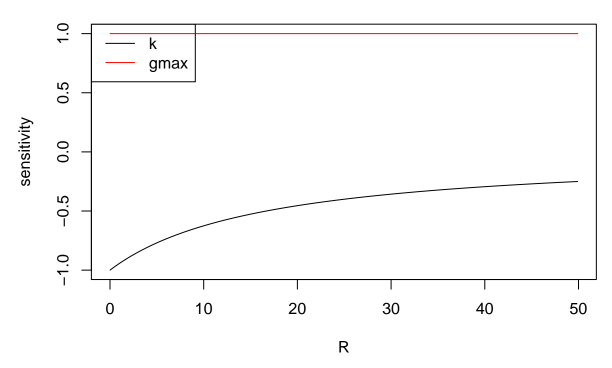
$$L_2(\Theta_j) = \sqrt{\sum S_{i,j}^2/n}$$

Try ?sensFun to identify the arguments required by sensFun (hint: in our case we will need 2+1 arguments). To do

- Store the ouput of sensFun in SF
- use summary, and plot to visulaize the sensitivity.

```
sF <- sensFun(f = Growth, parms= coef(Fita), R=seq(.01,50,.1))
summary(sF)
##
        value scale
                             L2
                                 Mean Min
                                                   N
                       L1
                                             Max
               16.7 0.46 0.022 -0.46
                                        -1 -0.25 500
## k
## gmax
          0.3
                0.3 1.00 0.045
                                 1.00
                                         1
                                            1.00 500
plot(sF)
```

#### All variables



A few question to reflect on:

- Why it the sensitivity to  $g_{max}$  always equal to 1?
- Why is the sensitivity to k always negative?
- Why is the sensitivity to k decreasing at large R values?

Note (by testing it) that  $L_1$  and  $L_2$  norms depend on the R value considered and on the initial parameter values.

## Distribution of parameters

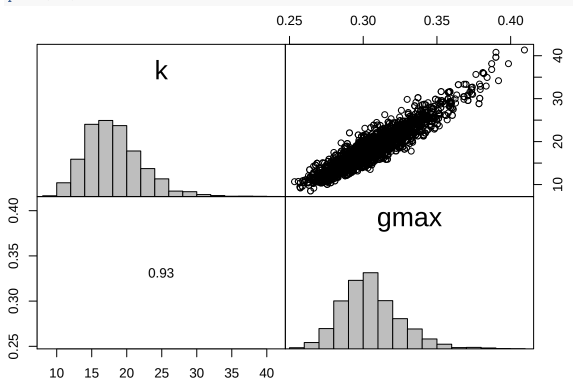
Instead of identifying a single set of "best" parameters, other approaches provides a probality distribution for parameter values.

It is better in the sense that is is statistically exploitable to derive error on model estimates. One such method is to use Monte Carlo Markov Chain. Here the transfer function returns an object ModCost, which is more complete than the simple residals we used previously.

```
ModelCostb <- function(P) {
  out <- Growth(P, datain$R)
  return(
   modCost(obs=datain,mod=out,x="R")</pre>
```

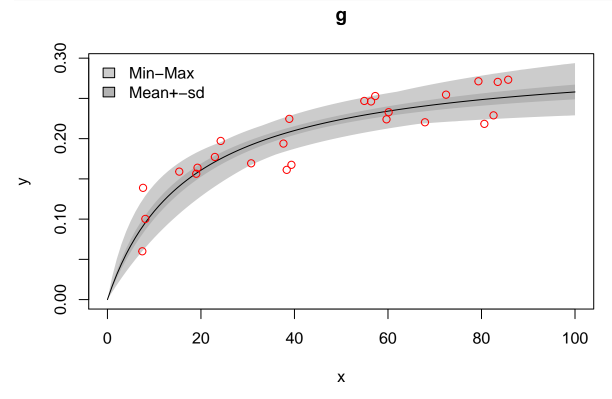
## number of accepted runs: 1597 out of 5000 (31.94%)

```
# This plots provides the selected parameter distributions pairs (MCa)
```



Given a distribution of parameters, such as provided from the previous procedure, the function sensRange allows to assess the distribution of model outputs. In this case we can consider it illustrates the uncertainties of model estimates deriving from the uncertainty on the calibrated parameters.

plot(summary(sR))
points(datain,col='red')



## References

Soetaert, Karline, and Thomas Petzoldt. 2010. "Inverse modelling, sensitivity and monte carlo analysis in R using package FME." Journal of Statistical Software 33 (3): 1-28.