Computer Lab 6 Computational Statistics

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Question 1: Genetic algorithm

In this assignment, you will try to perform one-dimensional maximization with the help of a genetic algorithm.

1. Define the function

$$f(x) := \frac{x^2}{e^x} - 2\exp(\frac{-9\sin x}{x^2 + x + 1})$$

```
# define the function
func = function(x){
  return((x^2/exp(x)) - 2 * exp(-(9 * sin(x))/ (x^2 +x +1)))
}
```

2. Define the function crossover()

for two scalars x and y it returns their "kid as (x+y)/2.

```
# crossover function
crossover = function(x,y){
  kid = (x+y)/2
  return(kid)
}
```

3. Define the function mutate()

that for a scalar x returns the result of the integer division $x2 \mod 30$. (Operation mod is denoted in R as %%).

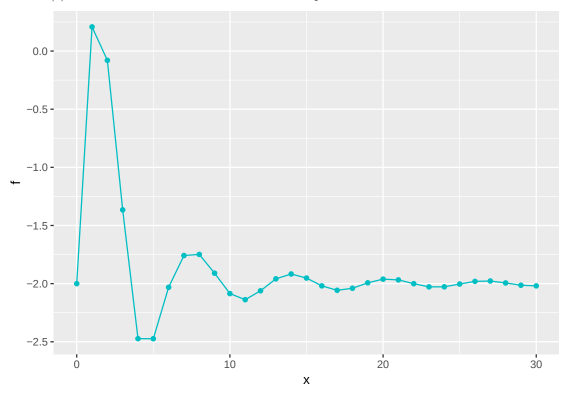
```
# mutate function
mutate = function(x){
  return(x^2 %%30)
}
```

4. Write a function

that depends on the parameters maxiter and mutprob and:

- (a) Plots function f in the range from 0 to 30. Do you see any maximum value?
- (b) Defines an initial population for the genetic algorithm as X = (0, 5, 10, 15, ..., 30).
- (c) Computes vector Values that contains the function values for each population point.
- (d) Performs maxiter iterations where at each iteration
 - i. Two indexes are randomly sampled from the current population, they are further used as parents (use sample()).
 - ii. One index with the smallest objective function is selected from the current population, the point is referred to as victim (use order()).
 - iii. Parents are used to produce a new kid by crossover. Mutate this kid with probability mutprob (use crossover(), mutate()).

- iv. The victim is replaced by the kid in the population and the vector *Values* is updated.
- v. The current maximal value of the objective function is saved.
- (e) Add the final observations to the current plot in another colour.



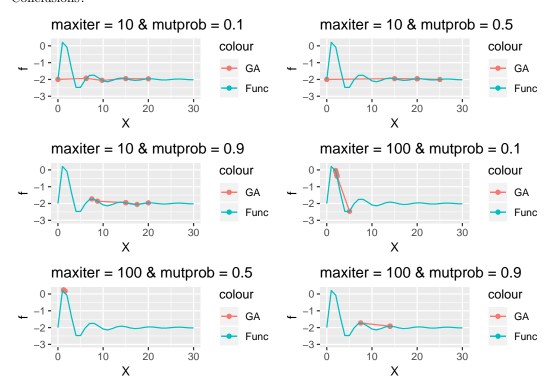
The maximum value might be between 0 and 2.

```
# initial population
X \leftarrow seq(0,30,5)
# the function values for each population points
Values <- func(X)
set.seed(1234567)
func4 <- function(pars, animation = F){</pre>
 maxiter = pars$maxiter
 mutprob = pars$mutprob
 name = pars$name
 tX <- X
 for(i in 1:maxiter) {
   samples <- sample(tX, 2, replace = F)</pre>
  id <- which.min(func(tX))</pre>
  kid <- crossover(samples[1],samples[2])</pre>
  if(runif(1)>mutprob){
    kid <- mutate(kid)</pre>
  }
  tX[id] <- kid
  tX <- sort(tX)
```

```
if(animation){  # Here we can see the change step by step
     plot(tX, func(tX), type = "b", xlim=c(0,30), ylim = c(-3,0.25), col="Blue")
     lines(x=seq(0,30),y=f(seq(0,30)))
     Sys.sleep(0.2)
   }
 }
dt <- data.frame(X=tX,f=func(tX))</pre>
 pl = ggplot(dt,aes(x=X,y=f,color="Blue"))+
   geom_point()+
   geom_line()+
   geom line(data=dataa,aes(x=x,y=f,color="Red"))+
   ylim(-3,0.25)+
   xlim(0,30) +
   ggtitle(name)+
   scale_color_discrete(labels=c("GA","Func"))
 return(pl)
}
```

5. Run your code with different combinations

of **maxiter**= 10, 100 and **mutprob**= 0.1, 0.5, 0.9. Observe the initial population and final population. Conclusions?



To begin with, the results with maxiter=100 have better average performance than the ones with maxiter=10. Most of them get the maximum around X=1, except for the one with mutprob=0.9.

Based on our code, mutprob represents threshold of mutation. If it is large, the children would be hard to

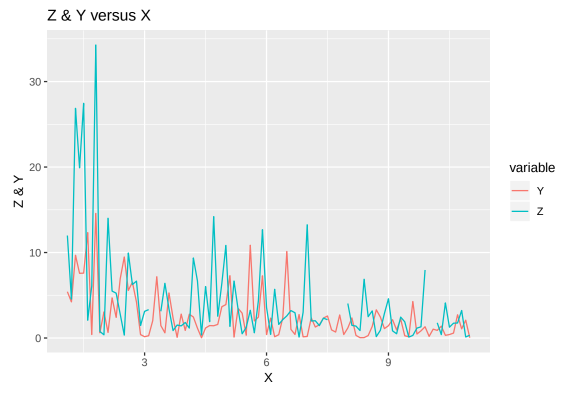
mutate. According to the plots, large mutprob might restrict the search range, which help us find a local optimal, not a global optimal.

Question 2: EM algorithm

The data file *physical.csv* describes a behavior of two related physical processes Y = Y(X) and Z = Z(X).

1. Make a time series plot

describing dependence of Z and Y versus X. Does it seem that two processes are related to each other? What can you say about the variation of the response values with respect to X?



variation versus X 10 -10 -20 -

It does not seem that the two processes are related to each other. In the beginning, the Z value does sleep off more than double of Y. Also, the movements of the processes rarely lie on top of each other. Based on the variation of such two responses, we can recognize the Z values are incomplete in some parts, the curve has gaps. However, the variation seems to decrease when X increases.

6 **X**

2. Derive an EM algorithm that estimates λ

3

Note that there are some missing values of Z in the data which implies problems in estimating models by maximum likelihood. Use the following model

$$Y_i \sim \exp(X_i/\lambda), \quad Z_i \sim \exp(X_i/2\lambda)$$

where λ is some unknown parameter. The goal is to derive an EM algorithm that estimates λ .

EM algorithm is a algorithm based on maximum likelihood estimation. We have to find the a probability density function (PDF) $p(x, z, \theta)$, where x is a sequence of observed data, z is latent data and θ are unknown parameters. Afterwards, there are two steps. Firstly, we expose the latent data z by maximum log-liklihood and current data x. Secondly, we get the estimate of parameters θ based on the maximum log-liklihood.

Expectation

In our case, we know the PDFs of Y and Z are

$$f(Y_i) = \frac{X_i}{\lambda} e^{-\frac{X_i}{\lambda} Y_i}, f(Z_i) = \frac{X_i}{2\lambda} e^{-\frac{X_i}{2\lambda} Z_i},$$

where the missing values in Z can be considered as latent data and the others are observed data. Since we consider that all the data are independent and we can thereby get the liklihood, which is

$$\mathcal{L}(\lambda|Y,Z) = \prod f(Y_i) \cdot \prod f(Z_i)$$

$$= \prod \left(\frac{X_i}{\lambda} e^{-\frac{X_i}{\lambda}Y_i}\right) \cdot \prod \left(\frac{X_i}{2\lambda} e^{-\frac{X_i}{2\lambda}Z_i}\right)$$

$$= \frac{\prod X_i}{\lambda^n} e^{-\frac{\sum X_i Y_i}{\lambda}} \cdot \frac{\prod X_i}{2^n \lambda^n} e^{-\frac{\sum X_i Z_i}{2\lambda}}.$$

Thus, the log-likelihood should be

$$\log \mathcal{L}(\lambda|Y,Z) = \prod (\ln X_i) - n \ln \lambda - \frac{\sum X_i Y_i}{\lambda} + \prod (\ln X_i) - n \ln(2\lambda) - \frac{1}{2\lambda} \sum X_i Z_i.$$

In the expectation step, we use maximum log-liklihood estimate to expose/express the missing data Z_{miss} by the other data (Y, Z_{obs}) in the last iteration, and thereby estimate our target parameter λ . Additionally, we replace the miss data with the expected value of Z.

$$\begin{aligned} Q(\lambda, \lambda_k) = & E[\log \mathcal{L}(\lambda|Y, Z_{miss}) | (\lambda_k, Y, Z_{obs})] \\ = & \prod (\ln X_i) - n \ln \lambda_k - \frac{\sum X_i Y_i}{\lambda_k} + \prod (\ln X_i) - n \ln(2\lambda_k) - \frac{1}{2\lambda_k} [\sum_{obs} X_i Z_i + \sum_{miss} X_i \frac{2\lambda_{k-1}}{X_i}] \\ = & 2 \prod (\ln X_i) - n \ln \lambda_k - \frac{\sum X_i Y_i}{\lambda_k} - n \ln(2\lambda_k) - \frac{1}{2\lambda_k} \sum_i X_i Z_i - \frac{1}{\lambda_k} \sum_i \lambda_{k-1}. \end{aligned}$$

maximization

To obtain estimate λ , we need to solve that the partial erivative of $Q(\lambda, \lambda_k)$ equals 0.

$$\nabla \lambda = -\frac{2n}{\lambda_k} + \frac{\sum X_i Y_i}{\lambda_k^2} + \frac{1}{2\lambda_k^2} \sum_{obs} X_i Z_i + \frac{1}{\lambda_k^2} \sum_{miss} \lambda_{k-1} = 0$$
$$\lambda_k = \frac{1}{4n} \left(2 \sum X_i Y_i + \sum_{obs} X_i Z_i + 2 \sum_{miss} \lambda_{k-1} \right)$$

3.Implement this algorithm in R

use $\lambda_0 = 100$ and convergence criterion "stop if the change in λ is less than 0.001". What is the optimal λ and how many iterations were required to compute it?

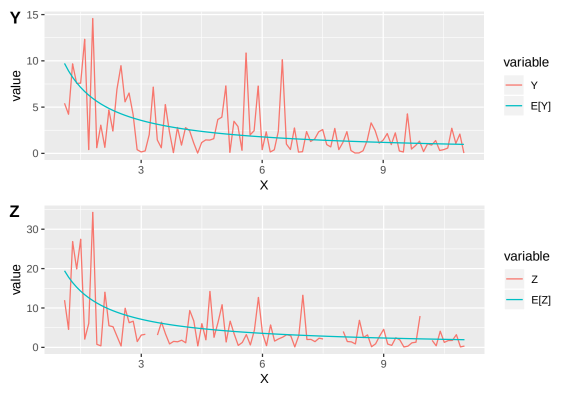
```
EM <- function(data2, eps, kmax=500, lda){
    X <- data2$X
    Y <- data2$Y
    Z <- data2$Z
    obs <- !is.na(Z)
    n <- length(X)
    m <- sum(is.na(Z))</pre>
```

```
# browser()
  loglik <- function(lda,ldap){</pre>
    return(2*sum(log(X))-n*log(lda)-(X%*%Y)/lda-n*log(2*lda)
           -(X[obs]%*%Z[obs])/(2*lda)-m*ldap/lda)
  }
  # initial state
  k <- 0
  llvalprev <- lda*10+10 # make some difference</pre>
  llvalcurr <- lda
  llk <- loglik(llvalcurr,llvalprev)</pre>
  print(c("iter","lda","llk"))
  print(c(k, llvalcurr,llk))
  while ((abs(llvalprev-llvalcurr)>eps) && (k<(kmax+1))){</pre>
    llvalprev <- llvalcurr</pre>
    # since we already know how to get argmax lda,
    # we can use the solution directly
    llvalcurr \leftarrow 1/(4*n)*(2*X%*%Y+X[obs]%*%Z[obs]+2*m*llvalprev)
    k<-k+1
    llk <- loglik(llvalcurr,llvalprev)</pre>
    print(c(k, llvalcurr,llk))
  }
  return(llvalcurr)
}
lda_opt <- EM(data2, eps=0.001, lda=100)</pre>
## [1] "iter" "lda" "llk"
## [1]
          0.0000 100.0000 -761.7647
          1.00000
## [1]
                   14.26782 -470.99635
## [1]
          2.00000
                    10.83853 -416.01655
## [1]
          3.00000
                   10.70136 -413.46921
## [1]
          4.00000
                   10.69587 -413.36664
## [1]
          5.00000
                     10.69566 -413.36253
```

According to the code, it is obvious that the log-liklihood increases during iterations. We get the optimal $\lambda = 10.69566$ after 5 iterations.

4. Plot E[Y] and E[Z] versus X

in the same plot as Y and Z versus X. Comment whether the computed λ seems to be reasonable.



In accordance with the plot, the paths (blue) of bath E[Y] and E[Z] pass through the original data (red), which show similar tendencies. We can therefore conclude that the λ we get is reasonable.

Appendix

```
knitr::opts_chunk$set(echo = TRUE, out.height = "280px")
# library used in this lab
library(ggplot2) # ex 2.1 - time series plot
library(gridExtra)
# clean the environment
rm(list=ls())
# define the function
func = function(x){
 return((x^2/\exp(x)) - 2 * \exp(-(9 * \sin(x))/(x^2 + x + 1)))
}
# crossover function
crossover = function(x,y){
 kid = (x+y)/2
 return(kid)
}
# mutate function
mutate = function(x){
 return(x^2 \\\\\30)
}
dataa \leftarrow data.frame(x=0:30,f=func(0:30))
plot1.4=ggplot(dataa,aes(x=x,y=f))+
 geom_line(color="#00BFC4")+
 geom_point(color="#00BFC4") # max x=1
plot1.4
# initial population
X \leftarrow seq(0,30,5)
# the function values for each population points
Values <- func(X)
set.seed(1234567)
func4 <- function(pars, animation = F){</pre>
 maxiter = pars$maxiter
 mutprob = pars$mutprob
 name = pars$name
 tX <- X
 for(i in 1:maxiter) {
   samples <- sample(tX, 2, replace = F)</pre>
   id <- which.min(func(tX))</pre>
   kid <- crossover(samples[1],samples[2])</pre>
   if(runif(1)>mutprob){
     kid <- mutate(kid)
   tX[id] <- kid
   tX <- sort(tX)
   if(animation){
                 # Here we can see the change step by step
     plot(tX, func(tX), type = "b", xlim=c(0,30), ylim = c(-3,0.25), col="Blue")
```

```
lines(x=seq(0,30), y=f(seq(0,30)))
      Sys.sleep(0.2)
    }
  }
dt <- data.frame(X=tX,f=func(tX))</pre>
  pl = ggplot(dt,aes(x=X,y=f,color="Blue"))+
    geom point()+
    geom line()+
    geom_line(data=dataa,aes(x=x,y=f,color="Red"))+
    ylim(-3,0.25)+
    xlim(0,30) +
    ggtitle(name)+
    scale_color_discrete(labels=c("GA","Func"))
  return(pl)
}
maxiter = c(10,100)
mutprob = c(0.1, 0.5, 0.9)
names = c("maxiter = 10 & mutprob = 0.1",
          "maxiter = 100 & mutprob = 0.5".
          "maxiter = 10 & mutprob = 0.9",
          "maxiter = 100 & mutprob = 0.1",
          "maxiter = 10 & mutprob = 0.5",
          "maxiter = 100 & mutprob = 0.9")
pairs = data.frame(maxiter=rep(maxiter,3),mutprob=rep(mutprob,2),name=names)
pairs = split(pairs,pairs[,3])
plot(arrangeGrob(grobs=lapply(t(pairs), func4)))
# clean the environment
rm(list=ls())
# load the data
data2 = read.csv("physical1.csv")
# Z & Y versus X
data21 <- reshape2::melt(data2, id.va = "X")</pre>
ggplot(data = data21,aes(x=X,y=value,color=variable)) +
  geom_line()+
  \# qeom\_line(aes(x = X, y=Y), color = "\#F8766D") +
  \# geom\_line(aes(x = X, y=Z), color = "\#00BFC4") +
  ggtitle("Z & Y versus X") +
  ylab("Z & Y")
ggplot(data = data.frame(X=data2\$X, variation=data2\$Y-data2\$Z), aes(x = X)) +
  geom_line(aes(y = variation)) +
  ggtitle("variation versus X")
EM <- function(data2, eps, kmax=500, lda){
  X <- data2$X
  Y <- data2$Y
  Z \leftarrow data2\$Z
  obs <- !is.na(Z)
  n <- length(X)
  m <- sum(is.na(Z))</pre>
```

```
# browser()
  loglik <- function(lda,ldap){</pre>
    return(2*sum(log(X))-n*log(lda)-(X%*%Y)/lda-n*log(2*lda)
           -(X[obs]%*%Z[obs])/(2*lda)-m*ldap/lda)
  # initial state
  k <- 0
  llvalprev <- lda*10+10 # make some difference</pre>
  llvalcurr <- lda
  llk <- loglik(llvalcurr,llvalprev)</pre>
  print(c("iter","lda","llk"))
  print(c(k, llvalcurr,llk))
  while ((abs(llvalprev-llvalcurr)>eps) && (k<(kmax+1))){</pre>
    llvalprev <- llvalcurr</pre>
    # since we already know how to get argmax lda,
    # we can use the solution directly
    llvalcurr \leftarrow 1/(4*n)*(2*X%*%Y+X[obs]%*%Z[obs]+2*m*llvalprev)
    k<-k+1
    llk <- loglik(llvalcurr,llvalprev)</pre>
    print(c(k, llvalcurr,llk))
  }
  return(llvalcurr)
}
lda_opt <- EM(data2, eps=0.001, lda=100)</pre>
X <- data2$X
EY <- c(lda_opt)/X
EZ <- 2*EY
data241 <- cbind(data2[,-3],data.frame(EY=EY))</pre>
data241 <- reshape2::melt(data241, id.va="X")</pre>
p1 <- ggplot(data241, aes(x=X, y=value, color=variable))+
  geom_line()+
  scale_color_discrete(labels=c("Y","E[Y]"))
data242 <- cbind(data2[,-2],data.frame(EY=EZ))</pre>
data242 <- reshape2::melt(data242, id.va="X")</pre>
p2 <- ggplot(data242, aes(x=X, y=value, color=variable))+</pre>
  geom_line()+
  scale_color_discrete(labels=c("Z","E[Z]"))
cowplot::plot_grid(p1, p2, labels = c('Y', 'Z'),ncol=1)
```