# Computer lab block 2

**Instructions**

* Create a report to the lab solutions in PDF.
* Be concise and do not include unnecessary printouts and figures produced by the software and not required in the assignments.
* **Include all your codes as an appendix into your report.**
* **Use set.seed(12345) for every piece of code that contains randomness**
* A typical lab report should 2-4 pages of text plus some amount of figures plus appendix with codes.
* The lab report should be submitted via LISAM before the deadline.

### Assignment 1. Ensemble methods

Your task is to learn some random forests using the function **randomForest** from the R package **randomForest**. The training data is produced by running the following R code:

x1<-runif(100)

x2<-runif(100)

trdata<-cbind(x1,x2)

y<-as.numeric(x1<x2)

trlabels<-as.factor(y)

The task is therefore classifying Y from X1 and X2, where Y is binary and X1 and X2 continuous. You should learn a random forest with 1, 10 and 100 trees, which you can do by setting the argument **ntree** to the appropriate value. Use **nodesize = 25** and **keep.forest = TRUE**. The latter saves the random forest learned. You need it because you should also compute the misclassification error in the following test dataset (use the function **predict** for this purpose):

set.seed(1234)

x1<-runif(1000)

x2<-runif(1000)

tedata<-cbind(x1,x2)

y<-as.numeric(x1<x2)

telabels<-as.factor(y)

plot(x1,x2,col=(y+1))

1. Repeat the procedure above for 1000 training datasets of size 100 and report the mean and variance of the misclassification errors. In other words, create 1000 training datasets of size 100, learn a random forest from each dataset, and compute the misclassification error in the **same** test dataset of size 1000. Report results for when the random forest has 1, 10 and 100 trees.
2. Repeat the exercise above but this time use the condition (x1<0.5) instead of (x1<x2) when producing the training **and** test datasets.
3. Repeat the exercise above but this time use the condition ((x1<0.5 & x2<0.5) | (x1>0.5 & x2>0.5)) instead of (x1<x2) when producing the training **and** test datasets. Unlike above, use **nodesize =** 12 for this exercise**.**
4. Answer the following questions:
   1. What happens with the mean and variance of the error rate when the number of trees in the random forest grows ?
   2. The third dataset represents a slightly more complicated classification problem than the first one. Still, you should get better performance for it when using sufficient trees in the random forest. Explain why you get better performance.
   3. Why is it desirable to have low error variance ?

### Assignment 2. Mixture models

Your task is to implement the EM algorithm for mixtures of multivariate Bernoulli distributions. Please use the R template below to solve the assignment. Then, use your implementation to show what happens when your mixture model has too few and too many components, i.e. set K=2,3,4 and compare results. Please provide a short explanation as well.

set.seed(1234567890)

max\_it <- 100 # max number of EM iterations

min\_change <- 0.1 # min change in log likelihood between two consecutive EM iterations

N=1000 # number of training points

D=10 # number of dimensions

x <- matrix(nrow=N, ncol=D) # training data

true\_pi <- vector(length = 3) # true mixing coefficients

true\_mu <- matrix(nrow=3, ncol=D) # true conditional distributions

true\_pi=c(1/3, 1/3, 1/3)

true\_mu[1,]=c(0.5,0.6,0.4,0.7,0.3,0.8,0.2,0.9,0.1,1)

true\_mu[2,]=c(0.5,0.4,0.6,0.3,0.7,0.2,0.8,0.1,0.9,0)

true\_mu[3,]=c(0.5,0.5,0.5,0.5,0.5,0.5,0.5,0.5,0.5,0.5)

plot(true\_mu[1,], type="o", col="blue", ylim=c(0,1))

points(true\_mu[2,], type="o", col="red")

points(true\_mu[3,], type="o", col="green")

# Producing the training data

for(n in 1:N) {

k <- sample(1:3,1,prob=true\_pi)

for(d in 1:D) {

x[n,d] <- rbinom(1,1,true\_mu[k,d])

}

}

K=3 # number of guessed components

z <- matrix(nrow=N, ncol=K) # fractional component assignments

pi <- vector(length = K) # mixing coefficients

mu <- matrix(nrow=K, ncol=D) # conditional distributions

llik <- vector(length = max\_it) # log likelihood of the EM iterations

# Random initialization of the paramters

pi <- runif(K,0.49,0.51)

pi <- pi / sum(pi)

for(k in 1:K) {

mu[k,] <- runif(D,0.49,0.51)

}

pi

mu

for(it in 1:max\_it) {

plot(mu[1,], type="o", col="blue", ylim=c(0,1))

points(mu[2,], type="o", col="red")

points(mu[3,], type="o", col="green")

#points(mu[4,], type="o", col="yellow")

Sys.sleep(0.5)

# E-step: Computation of the fractional component assignments

# Your code here

#Log likelihood computation.

# Your code here

cat("iteration: ", it, "log likelihood: ", llik[it], "\n")

flush.console()

# Stop if the lok likelihood has not changed significantly

# Your code here

#M-step: ML parameter estimation from the data and fractional component assignments

# Your code here

}

pi

mu

plot(llik[1:it], type="o")

### Assignment 3. High-dimensional methods

Data file **geneexp.csv** contains information about gene expression of three different cell types (column Cell Type). These cell types are CD4 and CD8 (two sorts of T cells) and CD19 (B cells). The aim of this assignment is to classify cells to the appropriate cell types using gene expressions and discover relevant genes for the given cell types.

1. Divide data into training and test sets (70/30) without scaling. Perform nearest shrunken centroid classification of training data in which the threshold is chosen by cross-validation. Provide a centroid plot and interpret it. How many genes were selected by the method? What meaning do positive and negative values have in the centroid plot? Can it happen that all values in the centroid plot are positive for some gene?
2. List the names of the 2 most contributing genes and find their alternative names in Google. Then, by checking this webpage <https://panglaodb.se/markers.html> find out whether these two genes are “marker genes” for given cell types. Report the test error of the model.
3. Compute the test error and the number of the contributing features for the following methods fitted to the training data:
   1. Elastic net with the binomial response and in which penalty is selected by the cross-validation
   2. Support vector machine with “vanilladot” kernel.

Compare the results of these models with the results of the nearest shrunken centroids (make a comparative table). Which model would you prefer and why?

1. Implement Benjamini-Hochberg method for the original data in which you test each cell type versus the remaining ones, and use t.test() for computing p-values. Present plots showing p-values and the rejection area for each cell type and interpret them. How many genes correspond to the rejected hypotheses for each cell type?

***Submission procedure***

***First read ‘Course Information.PDF’ at LISAM, folder ‘Course documents’***

**Assume that X is the current lab number, Y is your group number.**

**If you are neither speaker nor opponent for this lab,**

* Submit your report using *Lab X* item in the *Submissions* folder before the deadline.
* Make sure that you or some of your group members submits the group report using *Lab X group report*  in the *Submissions* folder before the deadline

**If you are a speaker for this lab,**

* Submit your report using *Lab X* item in the *Submissions* folder before the deadline.
* Make sure that you or some of your group members does the following before the deadline:
  + submits the group report using *Lab X group report*  in the *Submissions* folder before the deadline
  + Goes to *Sumissions* and opens item *Password X*. Then the student should put your group report into ZIP file *Lab X\_Group Y.zip* and protect it with a password you found in Password X
  + Uploads the file to *Collaborative workspace* 🡪*Lab X* folder

**If you are opponent for this lab,**

* Submit your report using *Lab X* item in the *Submissions* folder before the deadline.
* Make sure that you or some of your group members submits the group report using *Lab X group report*  in the *Submissions* folder before the deadline
* After the deadline for the lab has passed, go to Collaborative workspace🡪*Lab X* folder and download the appropriate ZIP file. Open the PDF in this ZIP file by using the password available in *Submission🡪Password X* item, read it carefully and prepare (in cooperation with other group members) **at least three questions/comments/improvement suggestions per lab assignment** in order to put them at the seminar.