Laboration report in Machine Learning

Computer lab 2 block 1 $_{732A99}$

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1 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)</pre>
```

The task is therefore classifying Y from X_1 and X_2 , where Y is binary and X_1 and X_2 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)</pre>
```

• 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.

```
# Function to create 1000 training datasets
sim_random_forest <- function(tedata, telabels, assignment, nodesize = 25) {
    misclass_ntree_1 <- c()
    misclass_ntree_10 <- c()
    misclass_ntree_100 <- c()

for(i in 1:1000) {
        x1 <- runif(100)
        x2 <- runif(100)

        # Assign y depending on assignment
        if(assignment == 1) {
            trdata <- cbind(x1, x2)
            y <- as.numeric(x1 < x2)
            trlabels <- as.factor(y)
        } else if(assignment == 2) {
            trdata <- cbind(x1, x2)</pre>
```

```
y \leftarrow as.numeric(x1 < 0.5)
       trlabels <- as.factor(y)</pre>
     } else if(assignment == 3) {
       trdata <- cbind(x1, x2)</pre>
       y \leftarrow as.numeric((x1 < 0.5 & x2 < 0.5) | (x1 > 0.5 & x2 > 0.5))
       trlabels <- as.factor(y)</pre>
     }
     # Train data to data.frame
     train <- data.frame(y = trlabels, x1, x2)</pre>
     # ntree = 1 ------
     r1 <- randomForest(y ~ ., data=train, ntree=1, nodesize=nodesize, keep.forest=TRUE)
     pred_r1 <- predict(r1, tedata)</pre>
     confusion_r1 <- table(telabels, pred_r1)</pre>
     misclass_ntree_1[i] <- (confusion_r1[1,2] + confusion_r1[2,1]) / sum(confusion_r1)
     # ntree = 10 ------
     r2 <- randomForest(y ~ ., data=train, ntree=10, nodesize=nodesize, keep.forest=TRUE)
     pred_r2 <- predict(r2, tedata)</pre>
     confusion_r2 <- table(telabels, pred_r2)</pre>
     misclass_ntree_10[i] <- (confusion_r2[1,2] + confusion_r2[2,1]) / sum(confusion_r2)
     # ntree = 100 -----
     r3 <- randomForest(y ~ ., data=train, ntree=100, nodesize=nodesize, keep.forest=TRUE)
     pred_r3 <- predict(r3, tedata)</pre>
     confusion_r3 <- table(telabels, pred_r3)</pre>
     misclass_ntree_100[i] <- (confusion_r3[1,2] + confusion_r3[2,1]) / sum(confusion_r3)
 }
 # Added random forest models to return
 return(list(misclass_ntree_1, misclass_ntree_10, misclass_ntree_100))
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)</pre>
y \leftarrow as.numeric(x1 < x2)
telabels <- as.factor(y)</pre>
ggplot(data.frame(x1,x2,y), aes(x1,x2,col = as.factor(y+1))) + geom_point() +
```

theme_bw() + labs(color = NULL)

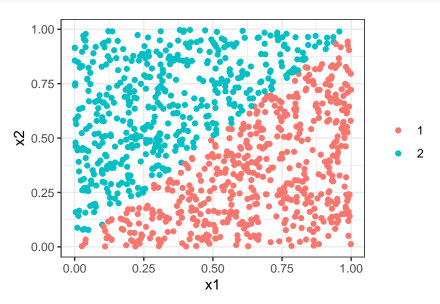


Figure 1: Testdata for first condition

Table 1: Mean and variance for first condition

	mean	variance
ntree = 1	0.2099	0.0034
ntree = 10	0.1350	0.0010
ntree = 100	0.1112	0.0009

• Repeat the exercise above but this time use the condition (x1 < 0.5) instead of (x1 < x2) when producing the training and test datasets.

```
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)
y <- as.numeric(x1 < 0.5)
telabels <- as.factor(y)

ggplot(data.frame(x1,x2,y), aes(x1,x2,col = as.factor(y+1))) + geom_point() +
    theme_bw() + labs(color = NULL)</pre>
```

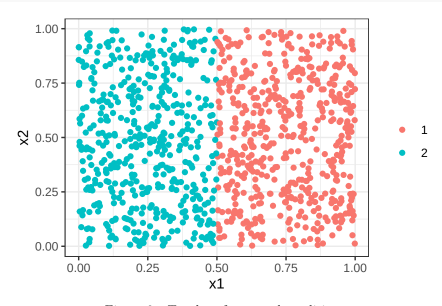


Figure 2: Testdata for second condition

Table 2: Mean and variance for second condition

	mean	variance
ntree = 1	0.0930	0.0173
ntree = 10	0.0138	0.0005
ntree = 100	0.0064	0.0001

• 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.

```
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)
y <- as.numeric((x1 < 0.5 & x2 < 0.5) | (x1 > 0.5 & x2 > 0.5))
telabels <- as.factor(y)

ggplot(data.frame(x1,x2,y), aes(x1,x2,col = as.factor(y+1))) + geom_point() +
theme_bw() + labs(color = NULL)</pre>
```

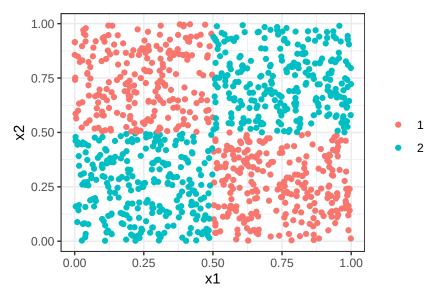


Figure 3: Testdata for third condition

Table 3: Mean and variance for third condition

	mean	variance
ntree = 1	0.2512	0.0133
ntree = 10	0.1228	0.0032
ntree = 100	0.0770	0.0014

- What happens with the mean error rate when the number of trees in the random forest grows? Why?
- 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.

Answer

The mean error decreases when the number of trees in the random forest grows. This happens because with more trees, the random forest becomes better at averaging out individual errors and capturing the pattern in the data. With more trees, random forest can fit a more complex boundary.

We obtain a lower mean for the third dataset compared to the first one, even though the third dataset represents a slightly more complicated classification problem than the first one for random forest. This occurs because the first dataset has a linear decision boundary that is not vertical or horizontal between the classes, and random forest is not a suitable model for capturing those decision boundarys. Other models, including simpler ones like logistic regression, might perform better in this case. Random forests are known for their ability to capture more complex relationships, which is why we observe better performance for the third dataset than the first dataset when the number of trees are larger.

2 Assignment 2. MIXTURE MODELS

Your task is to implement the EM algorithm for Bernoulli mixture model. 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 clusters, i.e. set M=2,3,4 and compare results. Please provide a short explanation as well. A Bernoulli mixture model is

$$p(\mathbf{x}) = \sum_{m=1}^{M} \pi_m Bern(\mathbf{x}|\mu_m)$$

where $\mathbf{x} = (x_1, ..., x_D)$ is a D-dimensional binary random vector, $\pi_m = p(y = m)$ and

$$Bern(\mathbf{x}|\mu_m) = \prod_{d=1}^{D} \mu_{m,d}^{x_d} (1 - \mu_{m,d})^{(1-x_d)}$$

where $\mu_m = (\mu_{m,1}, ..., \mu_{m,D})$ is a *D*-dimensional vector of probabilities. As usual, the log likelihood of the dataset $\{\mathbf{x}_i\}_{i=1}^n$ is

$$\sum_{i=1}^{n} \log p(\mathbf{x}_i)$$

Finally, in the EM algorithm, the parameter updates for the Bernoulli mixture model are:

$$\hat{\pi}_m = \frac{1}{n} \sum_{i=1}^n w_i(m)$$

$$\hat{\mu}_m = \frac{1}{\sum_{i=1}^n w_i(m)} \sum_{i=1}^n w_i(m) \mathbf{x}_i$$

where $w_i(m) = p(y_i = m|\mathbf{x}_i)$

Training data consists of 1 000 observations in 10 dimensions and are sampled as follows:

```
# Template code
set.seed(1234567890)
max_it <- 100 # max number of EM iterations
min_change <- 0.1 # min change in log lik between two consecutive 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)
```

```
# Producing the training data
for(i in 1:n) {
  m <- sample(1:3, 1, prob=true_pi)
  for(d in 1:D) {
    x[i,d] <- rbinom(1,1,true_mu[m,d])
  }
}</pre>
```

The true values for the mean, μ for each of the three clusters are presented in figure 4.

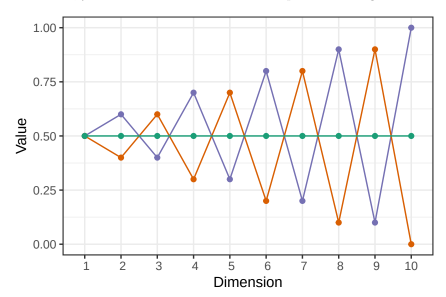


Figure 4: True value of μ .

2.1 EM-algorithm with 3 clusters

The EM-algorithm was implemented first for 3 clusters with the code as follows:

```
# Template code
# Creates empty variables for the EM-algorithm
M=3 # number of clusters
w <- matrix(nrow=n, ncol=M) # weights
pi <- vector(length = M) # mixing coefficients</pre>
mu <- matrix(nrow=M, ncol=D) # conditional distributions</pre>
llik <- vector(length = max_it) # log likelihood of the EM iterations</pre>
# Random initialization of the parameters
pi <- runif(M, 0.49, 0.51)
pi <- pi / sum(pi)
# Random initialization of mu for each cluster.
for(m in 1:M) {
  mu[m,] \leftarrow runif(D,0.49,0.51)
# End template code
bernoulli <- matrix(nrow=n, ncol=M)
for(it in 1:max_it) {
  \# E-step: Computation of the weights
  for (i in 1:1000){
    x i \leftarrow x[i,]
    bernoulli[i,1] \leftarrow prod(mu[1,]^x_i * (1-mu[1,])^(1-x_i))
    bernoulli[i,2] \leftarrow prod(mu[2,]^x_i * (1-mu[2,])^(1-x_i))
    bernoulli[i,3] \leftarrow prod(mu[3,]^x_i * (1-mu[3,])^(1-x_i))
    w[i,1] <- bernoulli[i,1] * pi[1] / sum(bernoulli[i,] * pi)</pre>
    w[i,2] <- bernoulli[i,2] * pi[2] / sum(bernoulli[i,] * pi)</pre>
    w[i,3] <- bernoulli[i,3] * pi[3] / sum(bernoulli[i,] * pi)
  }
  p_x <- bernoulli[,1]*pi[1] + bernoulli[,2]*pi[2] + bernoulli[,3]*pi[3]</pre>
  llik[it] <- sum(log(p_x))</pre>
  cat("iteration: ", it, "log likelihood: ", llik[it], "\n")
  # Stop if the log likelihood has not changed significantly
  if (it>1){
    if (abs(llik[it] - llik[it-1]) < min_change){</pre>
      # Saves a plot of the last iteration for mu
      plot_data <- data.frame(t(mu))</pre>
      p4 \leftarrow ggplot(plot_data, aes(x=1:10)) +
        geom_line(aes(y=X3), color="#7570B3") +
        geom point(aes(y=X3), color="#7570B3") +
```

```
geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom line(aes(y=X2), color="#1B9E77") +
      geom_point(aes(y=X2), color="#1B9E77") +
      theme bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
    # Exits the EM-algorithm
    break
 }
}
# M-step: ML parameter estimation from the data and weights
# Calculate new pi
pi <- 1/1000 * colSums(w)
# Calculates new mu
mu[1,] \leftarrow 1/sum(w[,1]) * colSums(w[,1] * x)
mu[2,] \leftarrow 1/sum(w[,2]) * colSums(w[,2] * x)
mu[3,] \leftarrow 1/sum(w[,3]) * colSums(w[,3] * x)
# Saves a plot of iteration 1 for mu
if(it == 1){
  plot_data <- data.frame(t(mu))</pre>
  p1 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X3), color="#7570B3") +
    geom_point(aes(y=X3), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X2), color="#1B9E77") +
    geom_point(aes(y=X2), color="#1B9E77") +
    theme_bw() +
    labs(x = "Dimension",
        y = "Value") +
    ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
# Saves a plot of iteration 9 for mu
if(it == 9){
  plot_data <- data.frame(t(mu))</pre>
  p2 \leftarrow ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X3), color="#7570B3") +
    geom_point(aes(y=X3), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X2), color="#1B9E77") +
```

```
geom_point(aes(y=X2), color="#1B9E77") +
      theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
     ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
  # Saves a plot of iteration 17 for mu
 if(it == 17){
   plot_data <- data.frame(t(mu))</pre>
   p3 <- ggplot(plot_data, aes(x=1:10)) +
      geom_line(aes(y=X3), color="#7570B3") +
      geom_point(aes(y=X3), color="#7570B3") +
      geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom_line(aes(y=X2), color="#1B9E77") +
      geom_point(aes(y=X2), color="#1B9E77") +
      theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
     ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
 }
}
## iteration: 1 log likelihood:
                                  -6931.482
## iteration: 2 log likelihood:
                                  -6929.074
## iteration: 3 log likelihood:
                                  -6928.081
## iteration: 4 log likelihood:
                                  -6920.57
## iteration: 5 log likelihood:
                                  -6868.29
```

```
## iteration: 6 log likelihood:
                                 -6646.505
## iteration: 7 log likelihood:
                                 -6403.476
## iteration: 8 log likelihood:
                                 -6357.743
## iteration: 9 log likelihood:
                                 -6351.637
## iteration: 10 log likelihood:
                                 -6349.59
## iteration: 11 log likelihood:
                                 -6348.513
## iteration: 12 log likelihood:
                                 -6347.809
## iteration: 13 log likelihood:
                                 -6347.284
## iteration: 14 log likelihood:
                                  -6346.861
## iteration: 15 log likelihood:
                                  -6346.506
## iteration: 16 log likelihood:
                                  -6346.2
## iteration: 17 log likelihood:
                                 -6345.934
## iteration: 18 log likelihood:
                                 -6345.699
## iteration: 19 log likelihood:
                                  -6345.492
## iteration: 20 log likelihood: -6345.309
## iteration: 21 log likelihood:
                                 -6345.147
## iteration: 22 log likelihood:
                                  -6345.003
## iteration: 23 log likelihood:
                                  -6344.875
## iteration: 24 log likelihood:
                                 -6344.762
```

iteration: 25 log likelihood: -6344.66
iteration: 26 log likelihood: -6344.57

The EM-algorithm converged after 26 iterations. In figure 5, the change in log likelihood is presented.

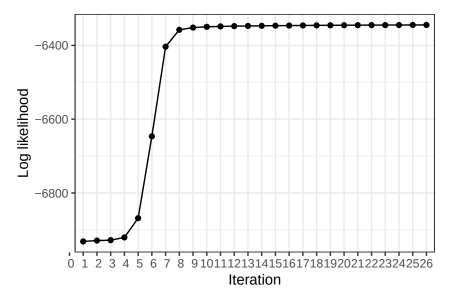


Figure 5: Change in log likelihood during the EM-algorithm.

In figure 6, the values of μ for iterations 1, 9, 17, 26 are presented for each cluster.

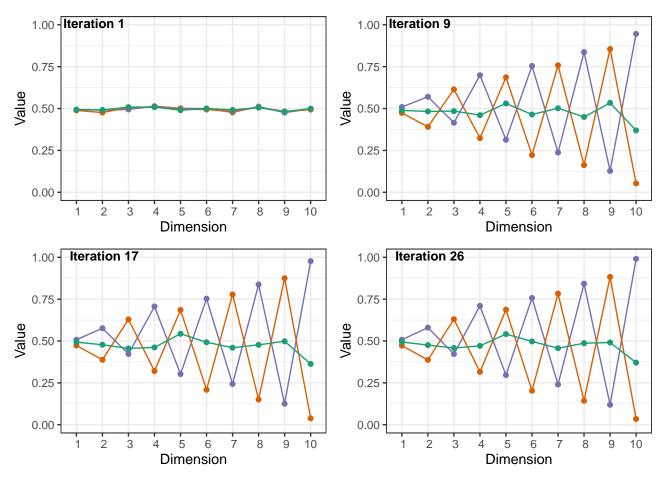


Figure 6: Different values for μ throughout the EM-algoritm with 3 clusters.

From figure 6, the EM-algorithm almost converged after 9 iterations to the three clusters. However, comparing the true values from figure 4, the algorithm does not converge exactly to the true values but still close.

2.2 EM-algorithm with 2 clusters

The EM-algorithm for 2 clusters used similar code as 3 clusters, however min change in log likelihood between two consecutive iterations was lowered from 0.1 to 0.05. Without the change, the algorithm converged after 3 iterations with both clusters still having values for μ around 0,5. The values for the log likelihood function are presented as follows:

```
## iteration:
               1 log likelihood:
                                   -6930.885
## iteration:
               2 log likelihood:
                                   -6929.223
               3 log likelihood:
                                   -6929.161
## iteration:
## iteration:
               4 log likelihood:
                                   -6928.816
               5 log likelihood:
                                   -6926.211
## iteration:
## iteration:
               6 log likelihood:
                                   -6906.763
## iteration:
               7 log likelihood:
                                   -6790.183
## iteration:
               8 log likelihood:
                                   -6499.677
  iteration:
               9 log likelihood:
                                   -6373.325
               10 log likelihood:
                                   -6364.13
## iteration:
               11 log likelihood:
                                    -6363.182
## iteration:
               12 log likelihood:
## iteration:
                                    -6362.96
## iteration:
               13 log likelihood:
                                   -6362.897
## iteration:
               14 log likelihood:
                                   -6362.878
```

The EM-algorithm converged after 14 iterations. In figure 7 the change in log likehood is presented.

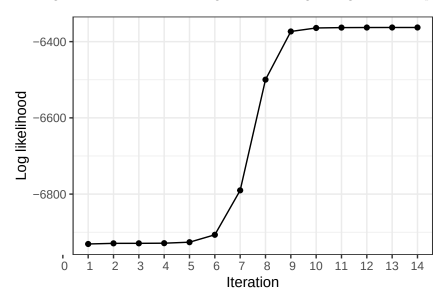


Figure 7: Change in log likelihood during the EM-algorithm.

In figure 8, the values of μ for iterations 1, 5, 10, 14 are presented for each cluster.

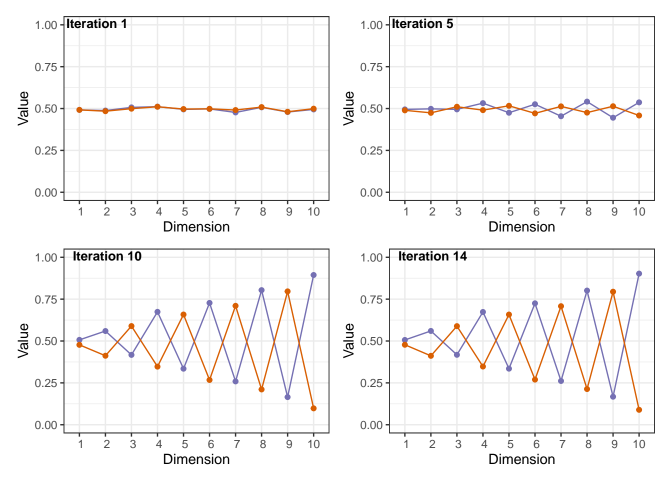


Figure 8: Different values for μ throughout the EM-algoritm with 2 clusters.

From figure 8, the EM-algorithm found the blue and red clusters but the green cluster could not be identified. Examining the probability matrix w a lot of observations had probabilities around 50% for each cluster and these are the observations that are from the green cluster.

2.3 EM-algorithm with 4 clusters

The code for EM-algorithm with 4 clusters had same stopping criterion as the template code and is presented under appendix. The values for the log likelihood function for each iterations of the EM-algorithm are as follows:

```
1 log likelihood:
                                   -6930.838
## iteration:
## iteration:
               2 log likelihood:
                                   -6928.641
## iteration:
               3 log likelihood:
                                   -6924.748
## iteration:
               4 log likelihood:
                                   -6896.25
## iteration:
               5 log likelihood:
                                   -6741.896
## iteration:
               6 log likelihood:
                                   -6452.658
## iteration:
               7 log likelihood:
                                   -6366.493
               8 log likelihood:
## iteration:
                                   -6359.764
## iteration:
               9 log likelihood:
                                   -6357.876
## iteration:
               10 log likelihood:
                                    -6356.372
## iteration:
               11 log likelihood:
                                    -6354.86
               12 log likelihood:
## iteration:
                                    -6353.31
               13 log likelihood:
## iteration:
                                    -6351.776
## iteration:
               14 log likelihood:
                                    -6350.33
               15 log likelihood:
                                    -6349.03
## iteration:
## iteration:
               16 log likelihood:
                                    -6347.908
## iteration:
               17 log likelihood:
                                    -6346.968
               18 log likelihood:
                                    -6346.196
## iteration:
## iteration:
               19 log likelihood:
                                    -6345.566
## iteration:
               20 log likelihood:
                                    -6345.055
                                    -6344.637
## iteration:
               21 log likelihood:
## iteration:
               22 log likelihood:
                                    -6344.293
## iteration:
               23 log likelihood:
                                    -6344.008
## iteration:
               24 log likelihood:
                                    -6343.768
## iteration:
               25 log likelihood:
                                    -6343.563
## iteration:
               26 log likelihood:
                                    -6343.387
## iteration:
               27 log likelihood:
                                    -6343.233
## iteration:
               28 log likelihood:
                                    -6343.097
               29 log likelihood:
## iteration:
                                    -6342.975
## iteration:
               30 log likelihood:
                                    -6342.864
## iteration:
               31 log likelihood:
                                    -6342.762
## iteration:
               32 log likelihood:
                                    -6342.668
```

The algorithm converged after 32 iterations and in figure 9 the loglikelihood are presented.

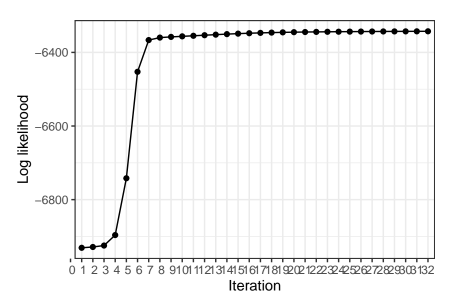


Figure 9: Change in log likelihood during the EM-algorithm.

The values for μ are presented for iterations 1, 12, 24, 32 in figure 10.

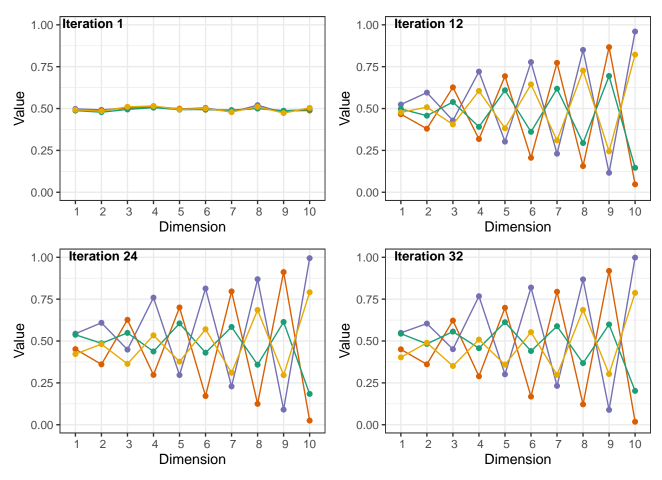


Figure 10: Different values for μ throughout the EM-algoritm with 4 clusters.

In figure 10, the mean values of the red and blue clusters are close to the true values in figure 4. Except for dimension 3, where the mean value is larger for the blue cluster. The green cluster in figure 4 was not identified, instead the cluster have been divided into two smaller clusters, green and yellow in figure 10.

3 Statement of Contribution

We worked on the assignment individually and later compared and discussed our solutions.

4 Appendix

The code used in this laboration report are summarised in the code as follows:

```
library(ggplot2)
library(tree)
library(knitr)
library(dplyr)
library(ggpubr)
library(randomForest)
knitr::opts_chunk$set(
  echo = TRUE,
  fig.width = 4.5,
  fig.height = 3)
x1 <- runif(100)
x2 <- runif(100)
trdata <- cbind(x1,x2)</pre>
y \leftarrow as.numeric(x1 < x2)
trlabels <- as.factor(y)</pre>
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1,x2)</pre>
y <- as.numeric(x1<x2)
telabels <- as.factor(y)</pre>
# Function to create 1000 training datasets
sim_random_forest <- function(tedata, telabels, assignment, nodesize = 25) {</pre>
  misclass_ntree_1 <- c()</pre>
  misclass_ntree_10 <- c()
  misclass_ntree_100 <- c()
  for(i in 1:1000) {
      x1 <- runif(100)
      x2 <- runif(100)
      # Assign y depending on assignment
      if(assignment == 1) {
```

```
trdata <- cbind(x1, x2)</pre>
       y \leftarrow as.numeric(x1 < x2)
       trlabels <- as.factor(y)</pre>
     } else if(assignment == 2) {
       trdata <- cbind(x1, x2)</pre>
       y \leftarrow as.numeric(x1 < 0.5)
       trlabels <- as.factor(y)</pre>
     } else if(assignment == 3) {
       trdata <- cbind(x1, x2)</pre>
       y \leftarrow as.numeric((x1 < 0.5 & x2 < 0.5) | (x1 > 0.5 & x2 > 0.5))
       trlabels <- as.factor(y)</pre>
     }
     # Train data to data.frame
     train <- data.frame(y = trlabels, x1, x2)</pre>
     # ntree = 1 ------
     r1 <- randomForest(y ~ ., data=train, ntree=1, nodesize=nodesize, keep.forest=TRUE)
     pred_r1 <- predict(r1, tedata)</pre>
     confusion_r1 <- table(telabels, pred_r1)</pre>
     misclass_ntree_1[i] <- (confusion_r1[1,2] + confusion_r1[2,1]) / sum(confusion_r1)
     # ntree = 10 -----
     r2 <- randomForest(y ~ ., data=train, ntree=10, nodesize=nodesize, keep.forest=TRUE)
     pred_r2 <- predict(r2, tedata)</pre>
     confusion_r2 <- table(telabels, pred_r2)</pre>
     misclass_ntree_10[i] <- (confusion_r2[1,2] + confusion_r2[2,1]) / sum(confusion_r2)
     # ntree = 100 -----
     r3 <- randomForest(y ~ ., data=train, ntree=100, nodesize=nodesize, keep.forest=TRUE)
     pred_r3 <- predict(r3, tedata)</pre>
     confusion_r3 <- table(telabels, pred_r3)</pre>
     misclass_ntree_100[i] \leftarrow (confusion_r3[1,2] + confusion_r3[2,1]) / sum(confusion_r3)
 }
 # Added random forest models to return
 return(list(misclass_ntree_1, misclass_ntree_10, misclass_ntree_100))
}
```

```
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)</pre>
y \leftarrow as.numeric(x1 < x2)
telabels <- as.factor(y)</pre>
theme_bw() + labs(color = NULL)
result1 <- sim_random_forest(tedata, telabels, assignment = 1)</pre>
r1 <- data.frame(mean = c(mean(result1[[1]]), mean(result1[[2]]), mean(result1[[3]])),
                variance = c(var(result1[[1]]), var(result1[[2]]), var(result1[[3]])))
rownames(r1) <- c("ntree = 1", "ntree = 10", "ntree = 100")
kable(r1, digits = 4, caption = "Mean and variance for first condition")
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)</pre>
y \leftarrow as.numeric(x1 < 0.5)
telabels <- as.factor(y)</pre>
ggplot(data.frame(x1,x2,y), aes(x1,x2,col = as.factor(y+1))) + geom_point() +
  theme_bw() + labs(color = NULL)
result2 <- sim_random_forest(tedata, telabels, assignment = 2)</pre>
r2 <- data.frame(mean = c(mean(result2[[1]]), mean(result2[[2]]), mean(result2[[3]])),
                 variance = c(var(result2[[1]]), var(result2[[2]]), var(result2[[3]])))
rownames(r2) <- c("ntree = 1", "ntree = 10", "ntree = 100")
kable(r2, digits = 4, caption = "Mean and variance for second condition")
```

```
set.seed(1234)
x1 <- runif(1000)
x2 <- runif(1000)
tedata <- cbind(x1, x2)</pre>
y \leftarrow as.numeric((x1 < 0.5 & x2 < 0.5) | (x1 > 0.5 & x2 > 0.5))
telabels <- as.factor(y)</pre>
ggplot(data.frame(x1,x2,y), aes(x1,x2,col = as.factor(y+1))) + geom_point() +
  theme_bw() + labs(color = NULL)
result3 <- sim_random_forest(tedata, telabels, assignment = 3, nodesize = 12)
r3 <- data.frame(mean = c(mean(result3[[1]]), mean(result3[[2]]), mean(result3[[3]])),
                 variance = c(var(result3[[1]]), var(result3[[2]]), var(result3[[3]])))
rownames(r3) <- c("ntree = 1", "ntree = 10", "ntree = 100")
kable(r3, digits = 4, caption = "Mean and variance for third condition")
# Template code
set.seed(1234567890)
max_it <- 100 # max number of EM iterations</pre>
min_change <- 0.1 # min change in log lik between two consecutive 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</pre>
true_mu <- matrix(nrow=3, ncol=D) # true conditional distributions</pre>
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)
# Producing the training data
for(i in 1:n) {
  m <- sample(1:3, 1, prob=true_pi)</pre>
  for(d in 1:D) {
    x[i,d] <- rbinom(1,1,true_mu[m,d])
  }
plot_data <- data.frame(t(true_mu))</pre>
```

```
ggplot(plot_data, aes(x=1:10)) +
  geom\_line(aes(y=X1), color="#7570B3") +
  geom_point(aes(y=X1), color="#7570B3") +
  geom_line(aes(y=X2), color="#D95F02") +
  geom_point(aes(y=X2), color="#D95F02") +
  geom_line(aes(y=X3), color="#1B9E77") +
  geom_point(aes(y=X3), color="#1B9E77") +
  theme bw() +
  labs(x = "Dimension",
       y = "Value") +
  scale_x_discrete(limits=c(1:10))
# Template code
# Creates empty variables for the EM-algorithm
M=3 # number of clusters
w <- matrix(nrow=n, ncol=M) # weights
pi <- vector(length = M) # mixing coefficients</pre>
mu <- matrix(nrow=M, ncol=D) # conditional distributions</pre>
llik <- vector(length = max_it) # log likelihood of the EM iterations</pre>
# Random initialization of the parameters
pi <- runif(M, 0.49, 0.51)
pi <- pi / sum(pi)
# Random initialization of mu for each cluster.
for(m in 1:M) {
  mu[m,] \leftarrow runif(D,0.49,0.51)
# End template code
bernoulli <- matrix(nrow=n, ncol=M)
for(it in 1:max_it) {
  # E-step: Computation of the weights
  for (i in 1:1000){
    x_i \leftarrow x[i,]
    bernoulli[i,1] <- prod(mu[1,]^x i * (1-mu[1,])^(1-x i))
    bernoulli[i,2] \leftarrow prod(mu[2,]^x_i * (1-mu[2,])^(1-x_i))
    bernoulli[i,3] \leftarrow prod(mu[3,]^x_i * (1-mu[3,])^(1-x_i))
    w[i,1] <- bernoulli[i,1] * pi[1] / sum(bernoulli[i,] * pi)</pre>
    w[i,2] <- bernoulli[i,2] * pi[2] / sum(bernoulli[i,] * pi)</pre>
    w[i,3] <- bernoulli[i,3] * pi[3] / sum(bernoulli[i,] * pi)
  }
  p_x <- bernoulli[,1]*pi[1] + bernoulli[,2]*pi[2] + bernoulli[,3]*pi[3]</pre>
  llik[it] <- sum(log(p_x))</pre>
  cat("iteration: ", it, "log likelihood: ", llik[it], "\n")
```

```
# Stop if the log likelihood has not changed significantly
if (it>1){
  if (abs(llik[it] - llik[it-1]) < min change){</pre>
    # Saves a plot of the last iteration for mu
    plot data <- data.frame(t(mu))</pre>
    p4 <- ggplot(plot_data, aes(x=1:10)) +
      geom_line(aes(y=X3), color="#7570B3") +
      geom_point(aes(y=X3), color="#7570B3") +
      geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom_line(aes(y=X2), color="#1B9E77") +
      geom_point(aes(y=X2), color="#1B9E77") +
      theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
    # Exits the EM-algorithm
    break
 }
}
# M-step: ML parameter estimation from the data and weights
# Calculate new pi
pi <- 1/1000 * colSums(w)
# Calculates new mu
mu[1,] \leftarrow 1/sum(w[,1]) * colSums(w[,1] * x)
mu[2,] \leftarrow 1/sum(w[,2]) * colSums(w[,2] * x)
mu[3,] \leftarrow 1/sum(w[,3]) * colSums(w[,3] * x)
# Saves a plot of iteration 1 for mu
if(it == 1){
  plot_data <- data.frame(t(mu))</pre>
  p1 <- ggplot(plot_data, aes(x=1:10)) +
    geom line(aes(y=X3), color="#7570B3") +
    geom_point(aes(y=X3), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X2), color="#1B9E77") +
    geom_point(aes(y=X2), color="#1B9E77") +
    theme_bw() +
    labs(x = "Dimension",
         y = "Value") +
    ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
}
```

```
# Saves a plot of iteration 9 for mu
  if(it == 9){
    plot_data <- data.frame(t(mu))</pre>
    p2 <- ggplot(plot_data, aes(x=1:10)) +
      geom_line(aes(y=X3), color="#7570B3") +
      geom_point(aes(y=X3), color="#7570B3") +
      geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom_line(aes(y=X2), color="#1B9E77") +
      geom_point(aes(y=X2), color="#1B9E77") +
      theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
  # Saves a plot of iteration 17 for mu
  if(it == 17){
    plot_data <- data.frame(t(mu))</pre>
    p3 <- ggplot(plot_data, aes(x=1:10)) +
      geom_line(aes(y=X3), color="#7570B3") +
      geom_point(aes(y=X3), color="#7570B3") +
      geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom_line(aes(y=X2), color="#1B9E77") +
      geom_point(aes(y=X2), color="#1B9E77") +
      theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
  }
plot_data <- data.frame(llik = llik[1:26])</pre>
ggplot(plot_data) +
  geom_point(aes(x=1:26, y=llik)) +
  geom_line(aes(x=1:26, y=llik)) +
 theme bw() +
  labs(x = "Iteration",
       y = "Log likelihood") +
  scale_x_discrete(limits=c(0:26))
ggarrange(p1, p2, p3, p4,
          labels=c("Iteration 1","Iteration 9","Iteration 17", "Iteration 26"),
          font.label = list(size = 10),
          hjust = -0.9,
          vjust = 2)
# EM-algorithm for 2 clusters
min_change <- 0.05 # min change in log lik between two consecutive iterations
M=2 # number of clusters
```

```
w <- matrix(nrow=n, ncol=M) # weights
pi <- vector(length = M) # mixing coefficients</pre>
mu <- matrix(nrow=M, ncol=D) # conditional distributions</pre>
llik <- vector(length = max_it) # log likelihood of the EM iterations</pre>
# Random initialization of the parameters
pi <- runif(M, 0.49, 0.51)
pi <- pi / sum(pi)
# Random initialization of mu for each cluster.
for(m in 1:M) {
  mu[m,] \leftarrow runif(D,0.49,0.51)
bernoulli <- matrix(nrow=n, ncol=M)
for(it in 1:max_it) {
  # E-step: Computation of the weights
  for (i in 1:1000){
    x_i \leftarrow x[i,]
    bernoulli[i,1] \leftarrow prod(mu[1,]^x_i * (1-mu[1,])^(1-x_i))
    bernoulli[i,2] \leftarrow prod(mu[2,]^x_i * (1-mu[2,])^(1-x_i))
    w[i,1] <- bernoulli[i,1] * pi[1] / sum(bernoulli[i,] * pi)</pre>
    w[i,2] <- bernoulli[i,2] * pi[2] / sum(bernoulli[i,] * pi)</pre>
  }
  p_x <- bernoulli[,1]*pi[1] + bernoulli[,2]*pi[2]</pre>
  llik[it] \leftarrow sum(log(p_x))
  cat("iteration: ", it, "log likelihood: ", llik[it], "\n")
  # Stop if the log likelihood has not changed significantly
  if (it>1){
    if (abs(llik[it] - llik[it-1]) < min_change){</pre>
      # Saves a plot of the last iteration for mu
      plot_data <- data.frame(t(mu))</pre>
      p8 <- ggplot(plot_data, aes(x=1:10)) +
        geom_line(aes(y=X2), color="#7570B3") +
        geom_point(aes(y=X2), color="#7570B3") +
        geom_line(aes(y=X1), color="#D95F02") +
        geom point(aes(y=X1), color="#D95F02") +
        theme bw() +
        labs(x = "Dimension",
             y = "Value") +
        ylim(0,1) +
        scale_x_discrete(limits=c(1:10))
      # Exits the EM-algorithm
```

```
break
 }
}
# M-step: ML parameter estimation from the data and weights
# Calculate new pi
pi <- 1/1000 * colSums(w)
# Calculates new mu
mu[1,] \leftarrow 1/sum(w[,1]) * colSums(w[,1] * x)
mu[2,] \leftarrow 1/sum(w[,2]) * colSums(w[,2] * x)
# Saves a plot of iteration 1 for mu
if(it == 1){
  plot_data <- data.frame(t(mu))</pre>
  p5 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
    geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    theme_bw() +
    labs(x = "Dimension",
         y = "Value") +
    ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
# Saves a plot of iteration 5 for mu
if(it == 5){
  plot_data <- data.frame(t(mu))</pre>
  p6 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
    geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    theme_bw() +
    labs(x = "Dimension",
         y = "Value") +
    ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
# Saves a plot of iteration 9 for mu
if(it == 10){
  plot_data <- data.frame(t(mu))</pre>
  p7 \leftarrow ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
    geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
```

```
theme_bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
  }
}
plot_data <- data.frame(llik = llik[1:14])</pre>
ggplot(plot_data) +
  geom_point(aes(x=1:14, y=1lik)) +
  geom_line(aes(x=1:14, y=llik)) +
  theme_bw() +
  labs(x = "Iteration",
       y = "Log likelihood") +
  scale_x_discrete(limits= c(0:14))
ggarrange(p5, p6, p7, p8,
          labels=c("Iteration 1","Iteration 5","Iteration 10", "Iteration 14"),
          font.label = list(size = 10),
          hjust = -0.9,
          vjust = 2)
min_change <- 0.1 # min change in log lik between two consecutive iterations
M=4 # number of clusters
w <- matrix(nrow=n, ncol=M) # weights
pi <- vector(length = M) # mixing coefficients</pre>
mu <- matrix(nrow=M, ncol=D) # conditional distributions</pre>
llik <- vector(length = max_it) # log likelihood of the EM iterations</pre>
\# Random initialization of the parameters
pi <- runif(M, 0.49, 0.51)
pi <- pi / sum(pi)
# Random initialization of mu for each cluster.
for(m in 1:M) {
  mu[m,] \leftarrow runif(D,0.49,0.51)
# End template code
bernoulli <- matrix(nrow=n, ncol=M)</pre>
for(it in 1:max_it) {
  # E-step: Computation of the weights
  for (i in 1:1000){
    x_i \leftarrow x[i,]
    bernoulli[i,1] \leftarrow prod(mu[1,]^x_i * (1-mu[1,])^(1-x_i))
    bernoulli[i,2] \leftarrow prod(mu[2,]^x_i * (1-mu[2,])^(1-x_i))
    bernoulli[i,3] \leftarrow prod(mu[3,]^x_i * (1-mu[3,])^(1-x_i))
    bernoulli[i,4] \leftarrow prod(mu[4,]^x_i * (1-mu[4,])^(1-x_i))
    w[i,1] <- bernoulli[i,1] * pi[1] / sum(bernoulli[i,] * pi)</pre>
```

```
w[i,2] <- bernoulli[i,2] * pi[2] / sum(bernoulli[i,] * pi)</pre>
  w[i,3] <- bernoulli[i,3] * pi[3] / sum(bernoulli[i,] * pi)</pre>
  w[i,4] <- bernoulli[i,4] * pi[4] / sum(bernoulli[i,] * pi)</pre>
p_x \leftarrow bernoulli[,1]*pi[1] + bernoulli[,2]*pi[2] + bernoulli[,3]*pi[3] + bernoulli[,4]*pi[4]
llik[it] <- sum(log(p_x))</pre>
cat("iteration: ", it, "log likelihood: ", llik[it], "\n")
# Stop if the log likelihood has not changed significantly
if (it>1){
  if (abs(llik[it] - llik[it-1]) < min_change){</pre>
    # Saves a plot of the last iteration for mu
    plot_data <- data.frame(t(mu))</pre>
    p12 <- ggplot(plot_data, aes(x=1:10)) +
      geom_line(aes(y=X2), color="#7570B3") +
      geom_point(aes(y=X2), color="#7570B3") +
      geom_line(aes(y=X1), color="#D95F02") +
      geom_point(aes(y=X1), color="#D95F02") +
      geom_line(aes(y=X3), color="#1B9E77") +
      geom_point(aes(y=X3), color="#1B9E77") +
      geom_line(aes(y=X4), color="#E6AB02") +
      geom_point(aes(y=X4), color="#E6AB02") +
      theme bw() +
      labs(x = "Dimension",
           y = "Value") +
      ylim(0,1) +
      scale_x_discrete(limits=c(1:10))
    # Exits the EM-algorithm
    break
 }
}
# M-step: ML parameter estimation from the data and weights
# Calculate new pi
pi <- 1/1000 * colSums(w)
# Calculates new mu
mu[1,] \leftarrow 1/sum(w[,1]) * colSums(w[,1] * x)
mu[2,] \leftarrow 1/sum(w[,2]) * colSums(w[,2] * x)
mu[3,] \leftarrow 1/sum(w[,3]) * colSums(w[,3] * x)
mu[4,] \leftarrow 1/sum(w[,4]) * colSums(w[,4] * x)
# Saves a plot of iteration 1 for mu
if(it == 1){
  plot_data <- data.frame(t(mu))</pre>
  p9 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
```

```
geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X3), color="#1B9E77") +
    geom_point(aes(y=X3), color="#1B9E77") +
    geom_line(aes(y=X4), color="#E6AB02") +
    geom_point(aes(y=X4), color="#E6AB02") +
   theme_bw() +
   labs(x = "Dimension",
         y = "Value") +
   ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
# Saves a plot of iteration 9 for mu
if(it == 12){
  plot_data <- data.frame(t(mu))</pre>
  p10 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
    geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X3), color="#1B9E77") +
    geom_point(aes(y=X3), color="#1B9E77") +
    geom_line(aes(y=X4), color="#E6AB02") +
    geom_point(aes(y=X4), color="#E6AB02") +
   theme_bw() +
    labs(x = "Dimension",
        y = "Value") +
   ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
# Saves a plot of iteration 17 for mu
if(it == 24){
  plot_data <- data.frame(t(mu))</pre>
  p11 <- ggplot(plot_data, aes(x=1:10)) +
    geom_line(aes(y=X2), color="#7570B3") +
    geom_point(aes(y=X2), color="#7570B3") +
    geom_line(aes(y=X1), color="#D95F02") +
    geom_point(aes(y=X1), color="#D95F02") +
    geom_line(aes(y=X3), color="#1B9E77") +
    geom_point(aes(y=X3), color="#1B9E77") +
    geom_line(aes(y=X4), color="#E6AB02") +
    geom_point(aes(y=X4), color="#E6AB02") +
    theme_bw() +
    labs(x = "Dimension",
         y = "Value") +
   ylim(0,1) +
    scale_x_discrete(limits=c(1:10))
```