report_lab1_block2

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1 Statement of Contribution

The contributions are distributed as follows:

Qingxuan Cui: Worked on assignment 1 and question 1 in assignment 4, with sharing and discussing the results.

Yanjie Lyu: Worked on assignment 2 (Mathematical derivation, and part of coding) and question 3 from assignment 4.

Yi Yang: Worked on assignment 2 (Part of coding and analysis) and question 2 from assignment 4.

After completing their respective assignments (including code writing and analysis), all results were shared and thoroughly discussed among the three members.

2 Assignment 1

2.1 Train Random Forest models with different hyperparameters and dataset

Node		Number of	Error	Error
Size	Condition	Trees	Mean	Variance
25	x1 <x2< td=""><td>1</td><td>0.109670</td><td>8.8e-04</td></x2<>	1	0.109670	8.8e-04
25	Ditto	10	0.109380	8.8e-04
25	Ditto	100	0.109230	8.7e-04
12	x1 < 0.5	1	0.006055	4.5e-05
12	Ditto	10	0.006074	4.7e-05
12	Ditto	100	0.006091	4.7e-05
12	(x1<0.5 & x2<0.5) (x1>0.5 & x2>0.5)	1	0.073000	1.3e-03
12	Ditto	10	0.072600	1.2e-03
12	Ditto	100	0.072500	1.2e-03

2.2 Answer for following questions

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

Among 3 different condition to generate label, the first one and the third one have the same trend that mean values decrease in their changes of mean classification error, with increase of node size, while the mean value for the second one increase slightly. I think the second one is a case, thus, we can conclude that, with node size increasing, the classification error decreases generally. As for reason, I think the prime one is that a bigger "forest" represents that data can be sampled for more times, subsequently we can acquire more base models trained by bootstrapped data sets, which are different after uniformly sampling. In that case, model can learn the feature from more different combinations of data subset, making the model less sensitive to outliers and noise.

2.2.2 The model trained with third dataset should get better performance than the first one. Why?

Certainly, the third model have better performance with much smaller mean classification error. The crux is that the third model set a smaller value about node size, in this case, the tree can split the data set for more times until the node size is less than node_size, which means the trees in third random forest model are deeper, and we can know that a deeper tree can catch more complicated relations between label and input data.

3 Assignment 2

3.1 Set M=2,3,4, the results show as follows

3.1.1 Set M=2,the results show as follows:

The number of iteration is : 12

The value of log likelihood: -6362.897

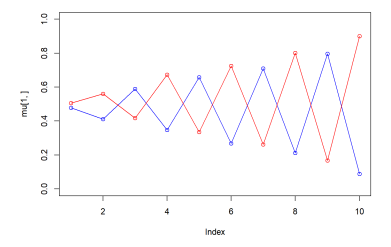


Figure 1: A.2.1: Value of mu after iterations with 2 clusters

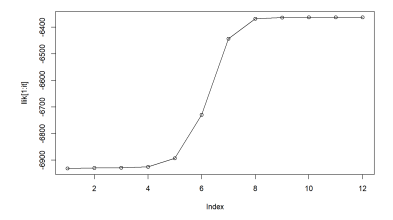


Figure 2: A.2.2: Value of log likelihood duiring iterations with 2 clusters

```
## The result of pi is:
## [1] 0.497125 0.502875
  The result of mu is :
                        [,2]
                                  [,3]
                                            [,4]
                                                       [,5]
                                                                 [,6]
             [,1]
                                                                            [,7]
## [1,] 0.4775488 0.4113939 0.5892308 0.3472420 0.6583712 0.2686589 0.7089490
## [2,] 0.5062860 0.5597531 0.4177551 0.6728856 0.3354854 0.7247188 0.2616231
             [,8]
                        [,9]
                                  [,10]
## [1,] 0.2118629 0.7957549 0.08905747
## [2,] 0.8007511 0.1678555 0.90027808
```

3.1.2 Set M=3, the results show as follows:

The number of iteration is : 26

The value of log likelihood: -6344.57

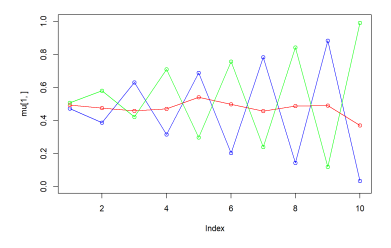


Figure 3: A.2.3: Value of mu after iterations with 3 clusters

```
## The result of pi is:
## [1] 0.3416794 0.2690298 0.3892909
## The result of mu is :
             [,1]
                       [,2]
                                  [,3]
                                            [,4]
                                                      [,5]
                                                                 [,6]
## [1,] 0.4727544 0.3869396 0.6302224 0.3156325 0.6875038 0.2030173 0.7832090
## [2,] 0.4939501 0.4757687 0.4584644 0.4711358 0.5413928 0.4976325 0.4569664
## [3,] 0.5075441 0.5800156 0.4221148 0.7100227 0.2965478 0.7571593 0.2400675
##
             [,8]
                       [,9]
                                  [,10]
## [1,] 0.1435650 0.8827796 0.03422816
## [2,] 0.4869015 0.4909904 0.37087402
## [3,] 0.8424441 0.1188864 0.99033611
```

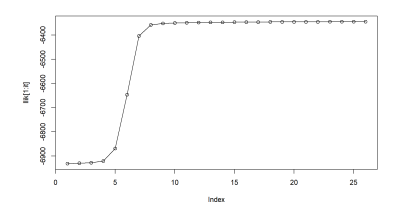


Figure 4: A.2.4: Value of log likelihood duiring iterations with 3 clusters

3.1.3 Set M=4, the results show as follows:

The number of iteration is : 44

The value of log likelihood: -6338.228

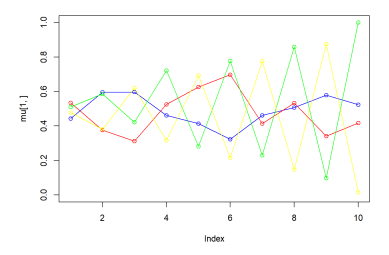


Figure 5: A.2.5: Value of mu after iterations with 4 clusters

The result of pi is:

[1] 0.1547196 0.1418652 0.3514089 0.3520062

The result of mu is :

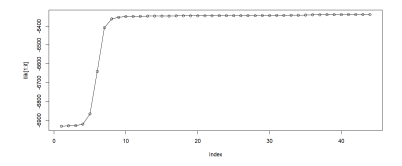


Figure 6: A.2.6: Value of log likelihood duiring iterations with 4 clusters

```
##
             [,1]
                        [,2]
                                  [,3]
                                            [,4]
                                                       [,5]
                                                                 [,6]
                                                                            [,7]
## [1,] 0.4426228 0.5955990 0.5973038 0.4611075 0.4148259 0.3224465 0.4616759
  [2,] 0.5347882 0.3763616 0.3116137 0.5256451 0.6254569 0.6980795 0.4139865
   [3,] 0.5103748 0.5869840 0.4219499 0.7218615 0.2825337 0.7763136 0.2299954
   [4,] 0.4781150 0.3812010 0.6195949 0.3165236 0.6926095 0.2166850 0.7756026
##
             [,8]
                         [,9]
##
                                   [,10]
## [1,] 0.5068223 0.57827821 0.52366273
  [2,] 0.5327794 0.34159869 0.41722943
  [3,] 0.8591562 0.09774851 0.99998228
## [4,] 0.1479707 0.87418437 0.01530099
```

3.2 Compare results

Case of M = 2: The log-likelihood value(-6362.897) is lower than M=3 and M=4, which shows that the model is relatively simple to capture the features of the data. The number of iterations(12) shows that it converges quickly but to a poor local optimal.

Case of M=3: The log-likelihood value (-6344.5) shows a significant improvement compared to M=2, which indicates the model fits the true distribution of the data more accurately. The number of iterations (26) shows a better balance between model complexity and model performance.

Case of M=4: The log-likelihood value (-6338.288) improves slightly compared to M=3, which means it does not show a significant improvement. The number of iterations (44) indicates increased model complexity complex to M=3, which may result in overfitting and the model may capture noise in the data.

Conclusion: A mixture model has too few clusters (e.g.,M=2 in this case) may lead to underfitting and can not capture necessary features from the data.On the contrary, a mixture with too many clusters(e.g.,M=4 in this case) may lead to excessive model complexity with little model performance. And clustering is not necessarily the case to minimize the "clustering loss", so we may perfer a smaller model(e.g.,M=3 in this case) over a large one even if the latter shows a slightly better log-likelihood.

4 Assignment 3 Theory

4.1 Impact of Ensemble Size (B) on Model Flexibility

My answer is no:

Flexibility means the model would be more complex to cope with some complicated data, which also means that it would be overfitted when B is big enough. (Bias tend to be smaller in train data and variance boom in test data) But actually we can see the effects after increasing B, based on the mean and variance:

$$\mathbb{E}\left[\frac{1}{B}\sum_{b=1}^{B} z_b\right] = \mu,$$

$$\operatorname{Var}\left[\frac{1}{B}\sum_{b=1}^{B} z_b\right] = \frac{1-\rho}{B}\sigma^2 + \rho\sigma^2.$$

Generally spkeaing, the variance would decrease while the bias remains. Summarize, there is no evidence representing the model would be flexible after B increasing.(P169)

4.2 The loss function used to train the boosted classifier at each iteration.

$$L(y \cdot f(x)) = \exp(-y \cdot f(x))$$

where

$$y \cdot f(x)$$

is the margin of the classifier. The ensemble members are added one at a time, and when member b is added, this is done to minimise the exponential loss of the entire ensemble. The reason why we choose the exponential loss is because it results in convenient closed form expressions. (page 177)

4.3 Use cross-validation to select the number of clusters

Data is split into K folds, and the model is trained on K-1 folds while leaving one fold out as a validation set. For each candidate value of M, we compute the average log-likelihood across all validation folds:

$$\frac{1}{K} \sum_{k=1}^{K} \log p \left(\{x_j'\}_{j=1}^{n_k} \mid \hat{\theta}_k, M \right),\,$$

The model with the largest average validation log-likelihood is then selected as the final clustering model with its optimal number of components M. (page 267)

5 Appendix

5.1 Code for assignment 1

```
generateTrainData = function(condition){
    x1 = runif(100)
    x2 = runif(100)
    trdata = cbind(x1,x2)
    y = as.numeric(eval(parse(text = condition)))
    trlabels = as.factor(y)
    return(cbind(trdata, trlabels))
}

generateTestData = function(condition){
```

```
set.seed(1234)
    x1 = runif(1000)
    x2 = runif(1000)
    tedata = cbind(x1,x2)
    y = as.numeric(eval(parse(text = condition)))
    telabels = as.factor(y)
    return(cbind(tedata, telabels))
library(randomForest)
trainFor3000Times = function(node_size, condition){
    test_data = generateTestData(condition)
    test_data[,3] = as.factor(test_data[,3])
    ntrees = c(1, 10, 100)
    rf_list = list()
    mes = vector()
    \# mean\_error\_list = list()
    # var_error_list = list()
    eval_list = list()
    eval = list()
    for (j in 1:3){
         for(i in 1:1000){
              train_data = generateTrainData(condition)
              rf = randomForest(as.factor(trlabels)~., data = train_data, ntrees = ntrees[j], nodesize = node_s
              output = predict(rf, newdata = test_data)
              me = mean(output != test_data[,3])
              mes = c(me, mes)
         }
         # report here
         trees = ntrees[j]
         mean_error = mean(mes)
         var_error = var(mes)
         eval$mean = mean_error
         eval$var = var_error
         eval_list[[as.character(trees)]] = eval
         \# mean_error_list[[as.character(trees)]] = mean_error
         cat(trees, " trees:\n")
         cat("Mean of misclassification error: ", mean_error, "\n")
         cat("Variance of misclassification error: ", var_error, "\n")
    }
    return(eval_list)
}
                                                          Condition: x1< x2 n")
cat("Node Size = 25
eval1 = trainFor3000Times(node_size = 25, condition = "x1<x2")
cat("Node Size = 12
                                                         Condition: x1<0.5\n")
eval2 = trainFor3000Times(node_size = 12, condition = "x1<0.5")
                                                         Condition: (x1<0.5 \& x2<0.5) | (x1>0.5 \& x2>0.5) | (x1>0.5 \& x2>
cat("Node Size = 12
eval3 = trainFor3000Times(node_size = 12, condition = "(x1<0.5 & x2<0.5)|(x1>0.5 & x2>0.5)")
```

5.2 code for assignment 2

```
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)
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(i in 1:n) {
  m <- sample(1:3,1,prob=true_pi)</pre>
 for(d in 1:D) {
    x[i,d] \leftarrow rbinom(1,1,true mu[m,d])
}
M=3 # number of clusters
#M=2
#M=4
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 \leftarrow runif(M, 0.49, 0.51)
pi <- pi / sum(pi)
for(m in 1:M) {
  mu[m,] \leftarrow runif(D,0.49,0.51)
}
рi
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 weights
  #Bern <- vector(length = n)
  #epsilon <- 1e-10
  #mu <- pmax(mu, epsilon)
```

```
#mu <- pmin(mu, 1 - epsilon)
 p_x <- numeric(length = n)</pre>
 for (i in 1:n) {
   pi_bern <- numeric(M)</pre>
   Bern <- rep(1, M)
   max_Bern <- 0</pre>
   for (m in 1:M){
     Bern[m] <- 1
     for (j in 1:D) {
       if(x[i,j]==1){
          Bern[m] <- Bern[m]*mu[m,j]</pre>
       }else{Bern[m] <- Bern[m]*(1-mu[m,j])}</pre>
     pi_bern[m] <- pi[m]*Bern[m]</pre>
     p_x[i] \leftarrow p_x[i] + pi_bern[m]
   for (m in 1:M) {
     w[i,m] <- pi_bern[m]/p_x[i]
   }
 }
 #Log likelihood computation.
 log_p_x \leftarrow log(p_x)
 llik[it] <- sum(log_p_x)</pre>
 cat("iteration: ", it, "log likelihood: ", llik[it], "\n")
 flush.console()
 # Stop if the lok likelihood has not changed significantly
 if(it > 1 && abs(llik[it] - llik[it - 1]) <= min_change){</pre>
   print(it)
   break
 }
 #M-step: ML parameter estimation from the data and weights
 for (m in 1:M) {
   N_m \leftarrow sum(w[,m])
  pi[m] <- N_m / n
  for (d in 1:D) {
     mu[m,d] \leftarrow sum(w[,m] * x[,d]) / N_m
}
```

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