Data Mining and Statistical Learning - Final Project

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I. QUESTION 1 – JACCARD SIMILARITY

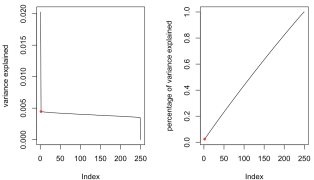
The definition of the Jaccard Similarity matrix is written as follow

$$Jac(A,B) = \frac{|A \cap B|}{|A \cup B|}$$

However, the way of calculating Jaccard Matrix is little different. THe way of computing the matrix is showed as follow

$$Jac(A,B) = \frac{|A = B = 1|}{|A = 1 \text{ or } B = 1|}$$

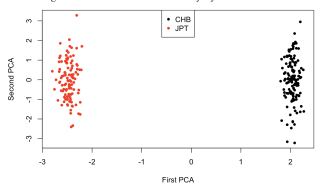
Fig. 1. Variance explained



The two figures show the variance explained and the cumulative variance explained by PCA. About 200 PCA can explain 80% of the variance.

However, as the plot below shows, the first two PCA can separate the two classes, Han Chinese in Beijing and Japanese in Tokyo pretty good.

Fig. 2. Classification of nationality by Jaccard Matrix



II. QUESTION 2 – CLASSIFICATION MODEL FOR CARDIAC ARRHYTHMIA

Methods can be applied:

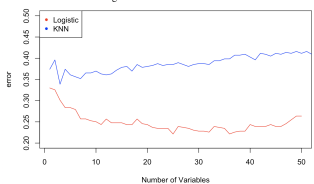
- 1. LDA
- 2. QDA
- 3. Best Subset
- 4. Forward Selection
- 5. Backward Selection
- 6. Stepwise Selection
- 7. Ridge Regression
- 8. Lasso regression
- 9. Elastic Net
- 10. Decision Tree
- 11. Bagging of the Trees
- 12. Random Forest
- 13. Gradient Boosting
- 14. ADA boosting
- 15. K-means
- 16. Hierarchical clustering

Methods not applied:

- 1. LDA is not used since it requires each covariate to be normally distributed.
- 2. Unsupervised learning is also not used since the label of the data is known.
- 3. Splines are not used also since most of the predictors are binary variables.

A. General Linear Model and KNN with variables selected by correlation

Fig. 3. GLM and KNN



The x lab of the figure is the number of the variables included in the model (selected by correlation).

As the figure shows, the Logistic regression has lower cross-validation error.

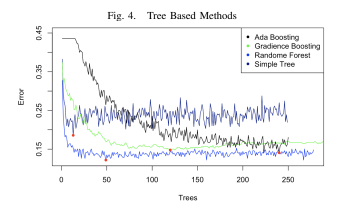
B. General Linear Model with variables selected by AIC

TABLE I
GENERAL LINEAR MODEL WITH VARIABLES SELECTED BY AIC

	Stepwise	Forward	Backward	subset
error	0.20	0.33	0.28	0.25

As the table shows, the stepwise selection has the lowest cv error.

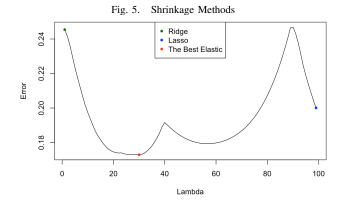
C. Tree Based Methods



The data was imputed at first, the column with missing value more than 80% was then deleted.

As the figure shows, the Random Forest model has the lowest cv error.

D. Shrinkage Methods



The Elastic net returns to the minimum cv error.

E. Model Comparison

Methods	error
logistic(Number of covariates = 38)	0.14
random forest (ntree = 122 , mtry = 49)	0.12
stepwise selection	0.20
elastic net ($\alpha = 0.2$, $\lambda = 29$)	0.17

The random Forest returns the best model with the lowest prediction error.

Compare to the paper. The prediction error is much smaller than the VFI5 method. However, the VFI5 method can categorize the result into 16 classes, it makes sense that VFI5 model has higher prediction error.

REFERENCES

- [1] Prokopenko D, Hecker J, Silverman EK, Pagano M, Nthen MM, Dina C, Lange C, Fier HL. Utilizing the Jaccard index to reveal population stratification in sequencing data: a simulation study and an application to the 1000 Genomes Project. Bioinformatics. 2016 May 1;32(9):1366-72. doi: 10.1093/bioinformatics/btv752. Epub 2015 Dec 31. PubMed PMID: 26722118; PubMed Central PMCID: PMC5860507.
- [2] H. Altay Guvenir, Burak Acar, Gulsen Demiroz, Ayhan Cekin "A Supervised Machine Learning Algorithm for Arrhythmia Analysis." Proceedings of the Computers in Cardiology Conference, Lund, Sweden, 1997.
- [3] https://www.genome.gov/10001688/international-hapmap-project/

APPENDIX

```
title: "Untitled"
output: html_document
```{r setup, include=FALSE}
knitr::opts_chunk$set(echo = TRUE)
library(MASS)
library(tidyverse)
library(nnet)
library(foreach)
library(doMC)
library(class)
library(glmnet)
library(caret)
library(tree)
library(randomForest)
library(pROC)
library(gbm)
library(xgboost)
library(xtable)
library(glmnet)
registerDoMC(10)
Q1
"\"{r}
dat.hapmap <- read.csv("/Users/hanchao/Library/Mobile_Documents/com~apple~CloudDocs/</pre>
Courses/Cornell_Semester_3/Statistical_Learning/FinalProject/SimulatedHapMap.csv")
, , ,
'''{r}
dat.exm <- dat.hapmap[,-1]</pre>
for (i in 1:ncol(dat.exm)) {
 if(!is.numeric(dat.exm[,i])){
 dat.exm[,i] <- as.numeric(as.character(dat.exm[,i]))</pre>
}
dat.exm[dat.exm > 0] <- 1
dat.exm <- as.data.frame(t(dat.exm))</pre>
t1 <- Sys.time()</pre>
smi <- list()</pre>
for (i in 1:ncol(dat.exm)) {
 a1 = dat.exm[,i] == dat.exm[,1:ncol(dat.exm)] & dat.exm[,i] != 0
 a2 = dat.exm[,i] != 0 | dat.exm[,1:ncol(dat.exm)] != 0
 smi[[i]] <- colSums(a1)/colSums(a2)</pre>
 print(i)
dat.jaccard <- do.call(cbind, smi)</pre>
dat.jaccard
```

```
t2 <- Sys.time()
+2-+1
dim(dat.jaccard)
. . .
Time difference of 9.851856 mins
Modify Chunk OptionsRun Current ChunkModify Chunk OptionsRun
All Chunks AboveRun Current ChunkModify Chunk OptionsRun All
Chunks AboveRun Current ChunkModify Chunk OptionsRun All Chunks
AboveRun Current Chunk
Console ~/
Console
Terminal
'''{r}
write.csv(dat.jaccard, "/Users/hanchao/Library/Mobile_Documents/
com~apple~CloudDocs/Courses/Cornell_Semester_3/Statistical_Learning/FinalProject/jaccard.csv")
. . .
'''{r}
dat.jaccard <- read.csv("/Users/hanchao/Library/Mobile_Documents/</pre>
com~apple~CloudDocs/Courses/Cornell_Semester_3/Statistical_Learning/FinalProject/jaccard.csv")
dat.jaccard <- dat.jaccard[,2:ncol(dat.jaccard)]</pre>
'''{r}
obj.prcomp <- prcomp(dat.jaccard, scale = TRUE)</pre>
'''{r}
par(mfrow = c(1,2))
plot((obj.prcomp$sdev^2)/sum(obj.prcomp$sdev^2), type = "1", ylab = "variance_explained")
points(x = 2, y = ((obj.prcomp\$sdev^2)/sum(obj.prcomp\$sdev^2))[2], col = "red", pch = 20)
plot(cumsum(obj.prcomp$sdev^2)/ sum(obj.prcomp$sdev^2) , type = "1",
ylab = "percentage_of_variance_explained")
points(x = 2, y = ((cumsum(obj.prcomp\$sdev^2) / sum(obj.prcomp\$sdev^2)))[2], col = "red", pch = 20)
, , ,
plot(-obj.prcomp$x[,1],obj.prcomp$x[,2], col = dat.hapmap$X, pch = 20, xlab = "First_PCA",
ylab = "Second PCA")
legend("top", y = as.character(unique(dat.hapmap$X)), col = c("black", "red"), pch = 20)
'''{r}
dat.ggplot <- data.frame(pca1 = obj.prcomp$x[,1],</pre>
pca2 = obj.prcomp$x[,2],
label = dat.hapmap$X)
Q 2
'''{r}
dat.arr <- read.table("/Users/hanchao/Library/Mobile_Documents/com~apple~CloudDocs/</pre>
Courses/Cornell_Semester_3/Statistical_Learning/FinalProject/arrhythmia.data", sep = ",", header = F)
names(dat.arr)[ncol(dat.arr)] <- "Y"</pre>
```

```
dat.arr\$Y \leftarrow ifelse(dat.arr\$Y > 1, yes = 1, no = 0)
, , ,
'''{r}
for (i in names(dat.arr)) {
 if(!is.numeric(dat.arr[,i])){
 print(i)
}
for (i in names(dat.arr)) {
 dat.arr[,i][dat.arr[,i] == "?"] <- NA</pre>
 dat.arr[,i] <- as.numeric(dat.arr[,i])</pre>
 if(max(dat.arr[,i], na.rm = T) == 16 \& min(dat.arr[,i], na.rm = T) == 1) {
 print(i)
}
for (i in names(dat.arr)) {
 if(mean(is.na(dat.arr[,i])) > 0.5){
 print(i)
}
#dat.arr <- dat.arr[,-which(names(dat.arr) == "V14")]</pre>
, , ,
LDA
'''{r}
cv.lda <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (i in 1:n) {
 train <- dat.splited[-i]
train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 data.frame(
 vairable = 1:279,
 cor = as.numeric(cor(dat.arr$Y, dat.arr[,-280]))
 arrange(desc(abs(cor))) -> dat.cor
 k.error <- NULL
 for (K in k) {
 fit.lda <- lda(Y~., data = train[, c(dat.cor$vairable[1:K], which(names(train) == y))]</pre>
 pred <- predict(fit.lda, newdata = valid)</pre>
 k.error[K] <- mean(pred$class != valid$Y)</pre>
 k.error
}
```

```
err.lda \leftarrow cv.lda(data = dat.arr, n = 5, k = 1:100, y = "Y")
. . .
'''{r}
cv.lda <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (K in k) {
 k.error <- NULL
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 data.frame(
 vairable = 1:279.
 cor = as.numeric(cor(dat.arr$Y, dat.arr[,-280]))
) 응>응
 arrange(desc(abs(cor))) -> dat.cor
 dat.train <- train[,c(dat.cor$vairable[1:K],280)]</pre>
 dat.valid <- valid[,c(dat.cor$vairable[1:K],280)]</pre>
 fit.lda <- lda(Y~., data = dat.train)</pre>
 pred <- predict(fit.lda, newdata = dat.valid)</pre>
 k.error[i] <- mean(pred$class != dat.valid$Y)</pre>
 knn.error[K] <- mean(k.error)</pre>
 z \leftarrow data.frame(k = k,
 error = knn.error)
 Z
}
err.lda <- cv.lda(data = dat.arr, n = 10, k = 1:50, y = "Y")
err.lda[which.min(err.lda$error),]
Logistic
'''{r}
cv.logistic <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 data.frame(
 vairable = 1:279,
 cor = as.numeric(cor(dat.arr$Y, dat.arr[,-280]))
) 응>응
 arrange(desc(abs(cor))) -> dat.cor
 k.error <- NULL
```

```
for (K in k) {
 fit.lda <- glm(Y~., data = train[, c(dat.cor$vairable[1:K],</pre>
 which(names(train) == y))] , family = binomial())
 pred <- predict(fit.lda, newdata = valid, type = "response")</pre>
 dat.roc <- roc(valid[,y]~pred)</pre>
 thres <- coords (dat.roc, "best") [1]
 pred <- ifelse(pred < thres, yes = 0, no = 1)</pre>
 k.error[K] <- mean(pred != valid$Y)</pre>
 k.error
}
err.logistic <- cv.logistic(data = dat.arr, n = 5, k = 1:100, y = "Y")
. . .
'''{r}
cv.multinomial <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (K in k) {
 k.error <- NULL
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 data.frame(
 vairable = 1:279,
 cor = as.numeric(cor(dat.arr$Y, dat.arr[,-280]))
) 응>응
 arrange(desc(abs(cor))) -> dat.cor
 dat.train <- train[,c(dat.cor$vairable[1:K],280)]</pre>
 dat.valid <- valid[,c(dat.cor$vairable[1:K],280)]</pre>
 fit.lda <- glm(Y~., data = dat.train, family = binomial)</pre>
 #fit.lda <- stepAIC(fit.lda)</pre>
 pred <- predict(fit.lda, newdata = dat.valid, type = "response")</pre>
 dat.roc <- roc(dat.valid$Y ~ pred)</pre>
 thres <-coords(dat.roc, "best")[1]</pre>
 pred <- ifelse(pred < thres, yes = 0, no = 1)</pre>
 k.error[i] <- mean(pred != dat.valid$Y)</pre>
 knn.error[K] <- mean(k.error)</pre>
 z \leftarrow data.frame(k = k,
 error = knn.error)
}
err.multinomial <- cv.multinomial(data = dat.arr, n = 5, k = 1:100, y = "Y")
err.multinomial[which.min(err.multinomial$error),]
. . .
KNN
```

```
'''{r}
cv.knn <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (K in k) {
 k.error <- NULL
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 Train <- data.frame(scale(train[,-which(names(train) == y)]),</pre>
 Class = train[,y])
 Valid <- data.frame(scale(valid[,-which(names(valid) == y)]),</pre>
 Class = valid[,y])
 pred.knn <- knn(Train[,-which(colnames(Train) == "Class")],</pre>
 Valid[,-which(colnames(Valid) == "Class")], Train[,"Class"], k = K)
 k.error[i] <- mean(pred.knn != Valid[, "Class"])</pre>
 knn.error[K] <- mean(k.error)</pre>
 z \leftarrow data.frame(k = k,
 knn.error)
 Z
}
dat.arr.knn <- dat.arr[,colMeans(is.na(dat.arr)) == 0]</pre>
for (i in names(dat.arr.knn)) {
 if (mean (duplicated (dat.arr.knn[,i])) > 0.9) {
 dat.arr.knn[,i] <- NA</pre>
}
dat.arr.knn <- dat.arr.knn[,colMeans(is.na(dat.arr.knn)) == 0]</pre>
dat.arr.knn <- data.frame(dat.arr.knn, Y = dat.arr$Y)</pre>
err.knn \leftarrow cv.knn(data = dat.arr.knn, n = 5, k = 1:100, y = "Y")
'''{r}
cv.qda <- function(data = data, n = "fold", k = "knn-numer", y = "Class"){</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 data.frame(
 vairable = 1:279,
 cor = as.numeric(cor(dat.arr$Y, dat.arr[,-280]))
) 응>응
 arrange(desc(abs(cor))) -> dat.cor
 }
 k.error <- NULL
 for (K in k) {
 fit.lda <- qda(Y~., data = train[, c(dat.cor$vairable[1:K], which(names(train) == y))]</pre>
 pred <- predict(fit.lda, newdata = valid)</pre>
 k.error[K] <- mean(pred$class != valid$Y)</pre>
```

```
k.error
}
. . .
Stepwise
'''{r}
cv.backward <- function(data = data, n = "fold", k = "knn-numer", y = "Class", direction = "both"){
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
 fit.lda <- glm(Y~., data = train, family = binomial())</pre>
 fit.lda <- step(fit.lda, glm(Y~., data = train, family = binomial()), direction = direction)</pre>
 pred <- predict(fit.lda, newdata = valid, type = "response")</pre>
 dat.roc <- roc(valid[,y]~pred)</pre>
 thres <- coords (dat.roc, "best") [1]
 pred <- ifelse(pred < thres, yes = 0, no = 1)</pre>
 k.error <- mean(pred != valid$Y)</pre>
 k.error
err.forward <- cv.backward(data = dat.arr.knn, n = 5, k = 1:100, y = "Y", direction = "forward")
err.stepwise <- cv.backward(data = dat.arr.knn, n = 5, k = 1:100, y = "Y", direction = "both")
err.backward <- cv.backward(data = dat.arr.knn, n = 5, k = 1:100, y = "Y")
tbl.aic <- data.frame(error = c(err.stepwise, err.forward,na.omit(err.backward)))</pre>
rownames(tbl.aic) <- c("Stepwise", "Forward", "Backward")</pre>
t(tbl.aic) %>%
 xtable()
plot(err.multinomial, type = "l", cex = 0.5, col = "red", ylim = c(0.2, 0.5),
xlab = "Number of Variables")
lines(err.knn, col = "blue")
legend(x="topleft", legend = c("Logistic", "KNN"), col = c("red", "blue"), pch = 20)
. . .
Tree
'''{r}
cv.tree2 <- function(data = data, n = "fold", k = "knn-numer", y = "Class") {</pre>
 knn.error <- NULL
 dd <- data[sample(nrow(data), nrow(data), replace = F),]</pre>
 dat.splited <- split(dd, rep(1:n, each=nrow(dd)/(n)))</pre>
 for (K in k) {
 k.error <- NULL
 for (i in 1:n) {
 train <- dat.splited[-i]</pre>
 train <- do.call(rbind, train)</pre>
 valid <- dat.splited[[i]]</pre>
```

```
my.tree <- tree(factor(Y)~., data = train)</pre>
 cv.model.pruned <- prune.misclass(my.tree, best=K)</pre>
 pred <- predict(cv.model.pruned, newdata = valid)[,2]</pre>
 dat.roc <- roc(valid$Y ~ pred)</pre>
 thres <- coords(dat.roc, "best")[1]
 pred.knn <- ifelse(pred < thres, yes = 0, no = 1)</pre>
 k.error[i] <- mean(pred.knn != valid[,"Y"])</pre>
 knn.error[K] <- mean(k.error)</pre>
 z \leftarrow data.frame(k = k,
 knn.error)
}
err.tree <- NULL
for (i in 2:500) {
 err.tree[i-1] <- as.numeric(na.omit(cv.tree2(data = dat.arr.tree, n = 5, k = i, y = "Y") $knn.error))
err.tree
plot(err.tree, type = "l")
points(which.min(err.tree), min(err.tree), pch = 20, col = "red")
'''{r}
dat.arr.tree <- dat.arr[,-which(names(dat.arr) == "V14")]</pre>
my.tree <- tree(factor(Y)~., data = dat.arr.tree)</pre>
mytree.cv <- cv.tree(my.tree,FUN=prune.misclass,K=10)</pre>
plot(x = mytree.cv$size, y = mytree.cv$dev, type = "b", xlab = "Size", ylab = "Deviance")
points(x = mytree.cv$size[which.min(mytree.cv$dev)],
y = mytree.cv$dev[which.min(mytree.cv$dev)], col = "red")
best.size <- mytree.cv$size[which(mytree.cv$dev==min(mytree.cv$dev))]</pre>
cv.model.pruned <- prune.misclass(my.tree, best=best.size[1])</pre>
plot(cv.model.pruned)
text(cv.model.pruned, use.n=TRUE, all=TRUE, cex=.8)
RF & bag of trees
hd.imputed <- rfImpute(Y~.-Y, data=dat.arr.tree)</pre>
err.rf1 <- list()
for (i in seq(1, 278, by = 1)) {
 obj.tree1 <- randomForest(factor(Y)~., data = hd.imputed, mtry=i,ntree=500)
 err.rf1[[i]] <- (obj.tree1$err.rate[,1])
a <- do.call(cbind, err.rf1)</pre>
err.rf <- apply(a, 2,min)
plot(na.omit(err.rf), type = "l", ylab = "Error", xlab = "Split",
main = "Random_Forest_and_Bagging_of_Tree")
```

```
points(which.min(err.rf), min(err.rf), pch = 20, col = "red")
GBM
'''{r}
library (Matrix)
##need to exclude missing
train.new <- na.omit(dat.arr.tree)</pre>
test.new <- na.omit(valid)</pre>
covariates_matrix = sparse.model.matrix(Y ~ ., data = train.new)[,-1]
output_vector = train.new[, 'Y'] == 1
#covariates_test_matrix = sparse.model.matrix(AHD ~ ., data = test.new)[,-1]
#output_test_vector = test.new[, 'AHD'] == "Yes"
##Create grid of parameters to try
xgb_grid = expand.grid(
 eta = c(0.1, 0.01, 0.001, 0.0001),
 max_depth = c(1,2)
for(i in 1:nrow(xgb_grid)){
assign(paste0("xgb_params.", i), list(
 objective = "binary:logistic",
 eta = xgb_grid[i,1],
 max.depth = xgb_grid[i,2],
 eval_metric = "error"
)) }
param.list <- list(xgb_params.1, xgb_params.2, xgb_params.3,</pre>
xgb_params.4, xgb_params.5, xgb_params.6,xgb_params.7,xgb_params.8)
set.seed(12345)
best.cv.error <- data.frame()</pre>
Eval.cv <- list()</pre>
for(i in 1:nrow(xgb_grid)){
xgb_cv = xgb.cv(params = param.list[[i]],
 data = covariates_matrix,
 label = output_vector,
 nrounds = 500,
 nfold = 5,
 prediction = TRUE,
 showsd = TRUE,
 stratified = TRUE,
 verbose = TRUE
)
Eval.cv[[i]] <- xgb_cv$evaluation_log[,c(1,4)]</pre>
best.cv.error[i,1] <- min(Eval.cv[[i]]$test_error_mean)</pre>
rownames(best.cv.error) <- c("eta=0.1,_depth=1", "eta=0.01,_depth=1", "eta=0.001,_depth=1", "eta=0.0001,
depth=1", "eta=0.1,_depth=2", "eta=0.01,_depth=2", "eta=0.001,_depth=2", "eta=0.0001,_depth=2")
which.min(best.cv.error$V1)
. . .
'''{r}
library('e1071')
##find shrinakge parameter
caretGrid.depth.1 <- expand.grid(interaction.depth=1, n.trees = 1000,</pre>
```

```
shrinkage=seg(0, 0.01, by = 0.001), n.minobsinnode=10)
set.seed(123)
gbm.caret.depth.1 <- caret::train(factor(Y) ~ ., distribution = "adaboost",</pre>
data = train.new, method = "gbm", trControl=trainControl(method="cv", number=5),
verbose=F, tuneGrid=caretGrid.depth.1)
best.tune.1 <- gbm.caret.depth.1$bestTune$shrinkage</pre>
names (gbm.caret.depth.1)
gbm.caret.depth.1$results
caretGrid.depth.2 <- expand.grid(interaction.depth=2, n.trees = 1:250,</pre>
shrinkage=seq(0, 0.01, by = 0.001), n.minobsinnode=10)
set.seed(123)
gbm.caret.depth.2 <- caret::train(factor(Y) ~ ., distribution = "adaboost",</pre>
data = train.new, method = "gbm", trControl=trainControl(method="cv", number=5),
verbose=F, tuneGrid=caretGrid.depth.2)
gbm.caret.depth.2$results[,c(1,2,4,5)]
best.tune.2 <- gbm.caret.depth.2$bestTune$shrinkage</pre>
##Fit the models with the parameters selected from cross validation
ada.train <- train.new
ada.train$AHD <- ifelse(ada.train$AHD=="No", 0, 1)</pre>
adaboost.depth.1 <- gbm(AHD ~ ., distribution="adaboost", data=ada.train,
shrinkage=best.tune.1, n.trees=2000)
adaboost.depth.2 <- gbm(AHD ~ ., distribution="adaboost", data=ada.train,</pre>
shrinkage=best.tune.2, n.trees=2000)
##Predict with 500 trees
pred.out.ada.depth.1 <- ifelse(predict(adaboost.depth.1, test.new, n.trees = 500,</pre>
type="response") >0.5, "Yes", "No")
misclass.ada.depth.1 <- mean(pred.out.ada.depth.1 != test.new$AHD)
misclass.ada.depth.1
, , ,
'''{r}
err.ada <- NULL
for (i in 1:250) {
 caretGrid.depth.2 <- expand.grid(interaction.depth=2, n.trees = i,</pre>
 shrinkage=seq(0, 0.01, by = 0.001), n.minobsinnode=10)
 gbm.caret.depth.2 <- caret::train(factor(Y) ~ ., distribution = "adaboost",</pre>
 data = train.new, method = "qbm", trControl=trainControl(method="cv", number=5),
 verbose=F, tuneGrid=caretGrid.depth.2)
 err.ada[i] <- gbm.caret.depth.2$results$Accuracy[which.max(gbm.caret.depth.2$results$Accuracy)]
}
. . .
'''{r}
plot(na.omit(err.rf), type = "l", ylab = "Error", xlab = "Trees", col = "blue",
ylim = c(0.12, 0.45), xlim = c(1, 240))
points(which.min(err.rf), min(err.rf), pch = 20, col = "red")
lines(err.tree, type = "l", col = "dark_blue")
points(which.min(err.tree), min(err.tree), pch = 20, col = "red")
lines(Eval.cv[[1]], col = "green")
points(which.min(Eval.cv[[1]]$test_error_mean), min(Eval.cv[[1]]$test_error_mean),
pch = 20, col = "red")
```

```
lines((1-err.ada))
points(x = which.min(1-err.ada), y = min((1-err.ada)), pch = 20, col = "red")
legend(x = "topright", legend = c("Ada_Boosting", "Gradience_Boosting", "Randome_Forest",
"Simple_Tree"), col = c("black", "green", "blue", "dark_blue"), pch = 20)
'''{r}
dat.arr.lasso <- na.omit(dat.arr.tree)</pre>
x \leftarrow model.matrix(Y^{\sim}.,dat.arr.lasso)[,-1] # First column corresponds to the intercept and is removed
y <- dat.arr.lasso$Y
1 < - seq(0, 1, by = 0.1)
err.shrinkage <- list()</pre>
for (i in 1:10) {
 enet.cv = cv.glmnet(x,y,alpha=l[i],nfolds=5)
 err.shrinkage[[i]] <- (enet.cv$cvm)</pre>
dat.lasso <- do.call(cbind,err.shrinkage)</pre>
dat.lasso[30,] == min(err.lasso)
err.lasso <- apply(dat.lasso, 1, min)</pre>
which(dat.lasso == min(err.lasso))
plot(err.lasso, type = "1", ylab = "Error", xlab = "Lambda")
points(x = 1, err.lasso[1], col = "dark_green", pch = 20)
points(x = length(err.lasso), err.lasso[length(err.lasso)], col = "blue", pch = 20)
points(x = which.min(err.lasso), min(err.lasso), col = "red", pch = 20)
legend(x = "top",legend = c("Ridge", "Lasso", "The_Best_Elastic"),
col = c("dark_green", "blue", "red"), pch = 20)
'''{r}
tbl.cp <- data.frame(error = c(min(err.logistic), min(err.rf), min(err.stepwise), min(err.lasso)))</pre>
rownames(tbl.cp) <- c("logistic", "random_forest", "stepwise", "Elastic")</pre>
(tbl.cp) %>%
 xtable()
which.min(err.rf)
```