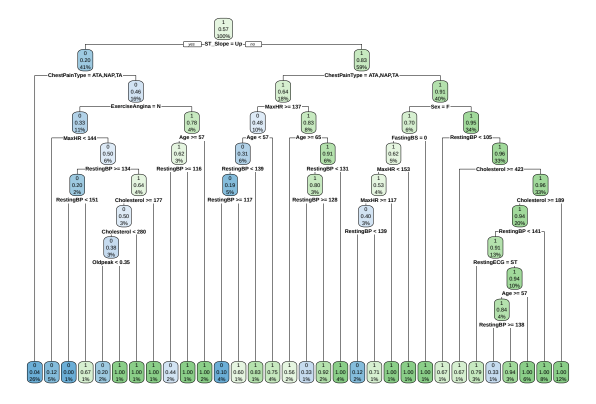
DS3 Hackathon Heart Disease Prediction

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Setup packages, load data, and further split the training (labeled) data into a subset for training and a subset for testing

```
library(tidyverse)
library(rpart)
library(rpart.plot)
library(randomForest)
library(gbm)
library(xgboost)
library(kableExtra)
library(ggplot2)
library(dplyr)
library(caret)
library(RCurl)
url_head <- "https://raw.githubusercontent.com/jeffreyz374/DS3-Hackathon-2023/main/heart/"
tr url <- paste0(url head, "heart train.csv")</pre>
heart_train <- getURL(tr_url)</pre>
heart_train <- read.csv(text = heart_train)</pre>
heart_train <- subset(heart_train, select = -c(X, id))
te_url <- paste0(url_head, "heart_test.csv")</pre>
heart test <- getURL(te url)</pre>
heart_test <- read.csv(text = heart_test)</pre>
heart_test <- subset(heart_test, select = -c(X))</pre>
heart_test_no_ids <- subset(heart_test, select = -c(id))</pre>
set.seed(1234)
train_idx <- sample(1:nrow(heart_train), round(0.8 * nrow(heart_train)))</pre>
train <- heart_train[train_idx,]</pre>
test <- heart_train[-train_idx,]</pre>
```

The first classifier we will use is the ordinary decision tree



• Since this tree is probably overfitting, we can prune the tree by looking at the complexity parameter:

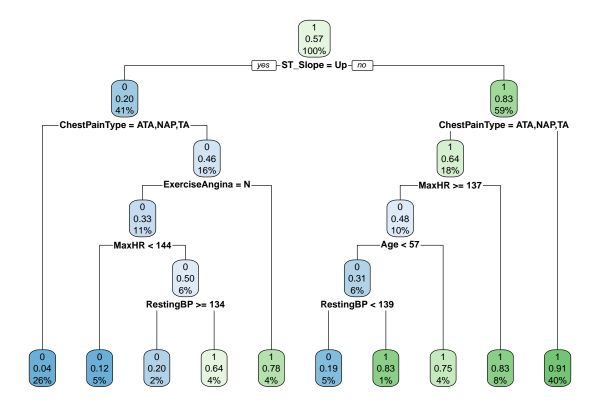
```
optimalcp <- heart_tree$cptable[which.min(heart_tree$cptable[,"xerror"]), "CP"]
optimalcp</pre>
```

[1] 0.00896861

• Prune the tree using the best value for the complexity parameter and draw the resulting tree:

```
heart_tree_prune <- prune(heart_tree, cp = optimalcp)

rpart.plot(heart_tree_prune)</pre>
```

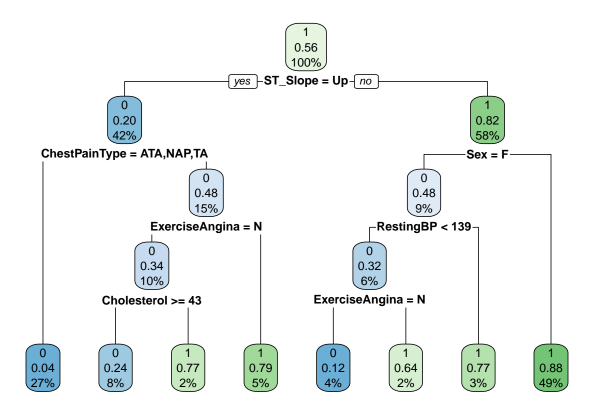


• Compute the test misclassification error

```
heart_pred <- predict(heart_tree_prune, test)</pre>
heart_pred <- as.data.frame(heart_pred)</pre>
heart_pred$HeartDisease <- ifelse(heart_pred$"0" > 0.5, 0, 1)
confmatrix_table <- table(true = test$HeartDisease, predicted = heart_pred$HeartDisease)</pre>
confmatrix_table
##
       predicted
##
   true
        0 1
##
      0 49 12
         9 59
##
misclass_err <- (confmatrix_table[1, 2] + confmatrix_table[2, 1]) / nrow(test)</pre>
misclass_err
```

[1] 0.1627907

• Fit the tree with the optimal complexity parameter to the full data:



• Predict on testing data:

• Join predictions with ids and export:

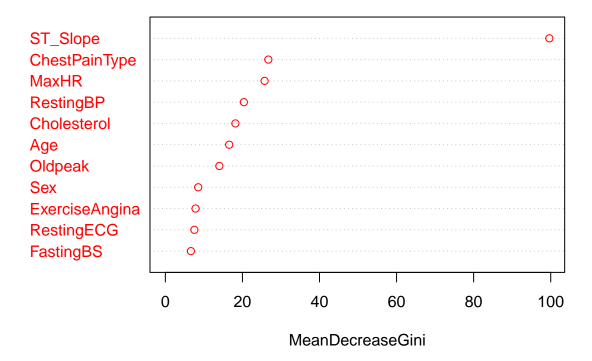
```
pruned_tree_predictions <- heart_pred_full$HeartDisease
x_pruned <- as.data.frame(list(id = heart_test$id, output = pruned_tree_predictions))
# Uncomment and change the destination directory to export
# write.csv(x_pruned, "~/Downloads/celestialSubmissionPruned.csv", row.names = FALSE)</pre>
```

• This classifier ultimately produced an accuracy of 0.8203108

The next classifier we will use is bagging

• We will set mtry equal to the number of features (all other parameters at their default values). We will also generate the variable importance plot using varImpPlot:

heart_bag



• Compute the test misclassification error:

```
heart_pred_bag <- predict(heart_bag, test)
heart_pred_bag <- as.data.frame(heart_pred_bag)
heart_pred_bag$HeartDisease <-
   ifelse(heart_pred_bag$heart_pred_bag == test$HeartDisease, 1, 0)

confmatrix_table <- table(true = test$HeartDisease,</pre>
```

```
predicted = heart_pred_bag$heart_pred_bag)

confmatrix_table

## predicted
## true 0 1
## 0 49 12
## 1 7 61

misclass_err <- (confmatrix_table[1, 2] + confmatrix_table[2, 1]) / nrow(test)
misclass_err</pre>
```

[1] 0.1472868

• This is a slight improvement from a single decision tree. Now, train it on the full data and predict on the testing data:

• Join predictions with ids and export:

```
x_bag <- as.data.frame(list(id = heart_test$id, output = heart_bag_pred_full$heart_bag_pred_full))
# Uncomment and change the destination directory to export
# write.csv(x_bag, "~/Downloads/heartSubmissionBag.csv", row.names = FALSE)</pre>
```

• This classifier ultimately produced an accuracy of 0.835001

The next classifier we will use is random forest

• This time, we will use randomForest with the default parameters. We will again generate the variable importance plot using varImpPlot:

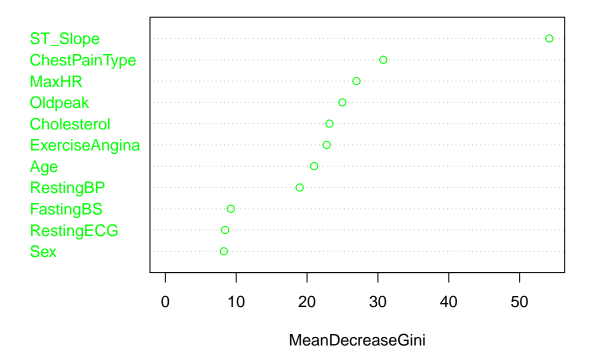
```
heart_rf <- randomForest(as.factor(HeartDisease)~., data = train, na.action = na.omit)

# Error rate
sum(heart_rf$err.rate[,1])

## [1] 66.15847

# Variable importance plot
varImpPlot(heart_rf, n.var = 11, col = "green")</pre>
```

heart_rf



• Compute the test misclassification error:

```
## predicted
## true 0 1
## 0 51 10
## 1 3 65

misclass_err <- (confmatrix_table[1, 2] + confmatrix_table[2, 1]) / nrow(test)
misclass_err</pre>
```

[1] 0.1007752

• This is an improvement from bagging. Now, train it on the full data and predict on the test data:

• Join predictions with ids and export:

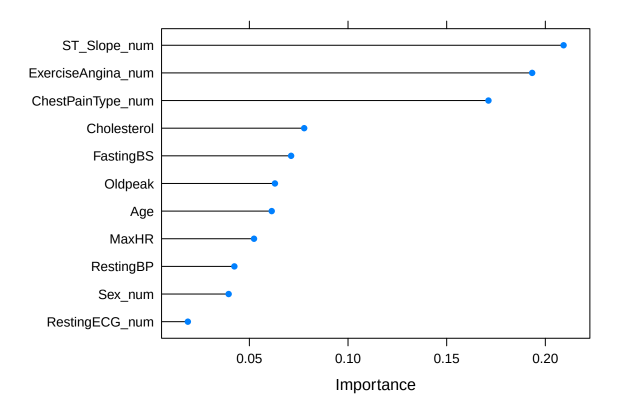
• This classifier ultimately produced an accuracy of 0.8530977

The final classifier we will use is extreme gradient boosting

• Clean the dataset to fit requirements necessary for extreme gradient boosting:

```
set.seed(1234)
# Replace categorical variables with numerical placeholders
heart_train$Sex_num <- ifelse(heart_train$Sex=="M",1,0)
heart_train$ChestPainType_num <-
  ifelse(heart_train$ChestPainType=="ATA", 1,
         ifelse(heart_train$ChestPainType=="NAP", 2,
                ifelse(heart_train$ChestPainType=="ASY",3,0)))
heart_train$RestingECG_num <-
  ifelse(heart_train$RestingECG=="Normal", 1,
         ifelse(heart_train$RestingECG=="ST",2,0))
heart_train$ExerciseAngina_num <- ifelse(heart_train$ExerciseAngina=="Y", 1, 0)
heart_train$ST_Slope_num <-
  ifelse(heart train$ST Slope=="Up", 1, ifelse(heart train$ST Slope=="Flat", 2, 0))
# Reassign labeled data, now with newly created numerical variables
heart_train <- heart_train %>%
  select(-c(Sex, ChestPainType, RestingECG, ExerciseAngina, ST_Slope))
# Re-split the labeled data into training and testing datasets
train <- sample(1:nrow(heart_train), floor(nrow(heart_train) * 0.8))</pre>
test <- setdiff(1:nrow(heart_train), train)</pre>
```

• Train extreme gradient boosting model with xgboost and perform a grid search for tuning the number of trees and the maximum depth of the tree. Then, we perform 10-fold cross-validation and determine the variable importance:



• Compute the test MSE:

```
yhat_xgb <- predict(heart_xgb, newdata = heart_train[test,])
mean((yhat_xgb - heart_train[test, "HeartDisease"]) ** 2)</pre>
```

[1] 0.09101545

• This is an improvement from random forest. Now, train it on the full data, and predict on the test data:

• Join predictions with ids and export:

• This classifier ultimately produced an accuracy of 0.8595912

Conclusion

All in all, the accuracies we received can be summarized in the below table:

```
accs <- c("0.8203108", "0.835001", "0.8530977", "0.8595912")
rownames <- c("Pruned Decision Tree", "Bagging", "Random Forest", "Extreme Gradient Boosting")
colname <- c("Accuracy")
acc_table <- cbind(accs)
rownames(acc_table) <- rownames
colnames(acc_table) <- colname
acc_table %>%
   kable(align = c("r"))
```

	Accuracy
Pruned Decision Tree	0.8203108
Bagging	0.835001
Random Forest	0.8530977
Extreme Gradient Boosting	0.8595912