# ISLR | Chapter 8 Exercises

Marshall McQuillen
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## Conceptual

### 1

(sketch on following page)

### $\mathbf{2}$

Consider the regression setting; when each tree has depth one (meaning there is only one split), each tree will only consist of one split on one attribute, and the predicted response for those two groups will be the mean response (of the training data) of those two groups. Therefore, the base model will take the form  $f_j(x_j) = \beta_j I(x_{ji} \le t) + \beta'_j I(x_{ji} > t)$ , where t is the value in the range of  $x_j$  that dictates the split (in order to minimize the RSS). Knowing this, one can walk through algorithm 8.2, as shown below:

$$f(\hat{x}) = 0 \tag{1}$$

$$f_1(x_1) = \beta_1 I(x_{1i} \le t) + \beta_1' I(x_{1i} > t)$$
(2)

$$\hat{f(x)} = \hat{f(x)} + \lambda f_1(x_1) \tag{3}$$

$$r_i = r_i - \lambda f_1(x_i) \tag{4}$$

$$f_2(x_2) = \beta_2 I(x_{2i} \le t) + \beta_2' I(x_{2i} > t)$$
(5)

$$\hat{f(x)} = \hat{f(x)} + \lambda f_2(x_2) \tag{6}$$

$$r_i = r_i - \lambda f_2(x_i) \tag{7}$$

$$\vdots$$
 (8)

$$f_i(x_i) = \beta_i I(x_{ii} \le t) + \beta_i' I(x_{ii} > t) \tag{9}$$

$$\hat{f(x)} = \hat{f(x)} + \lambda f_i(x_i) \tag{10}$$

$$r_i = r_i - \lambda f_i(x_i) \tag{11}$$

$$\vdots (12)$$

$$f(x) = \sum_{j=1}^{P} f_j(X_j) \quad where \quad f_j(X_j) = \beta_j I(x_{ji} \le t) + \beta'_j I(x_{ji} > t)$$
 (13)

(14)

Note that  $\beta_j$  is the mean of the response for all values of  $x_j$  that are less than or equal to t, and  $\beta'_j$  is the mean response for all values of  $x_j$  that are greater than t.

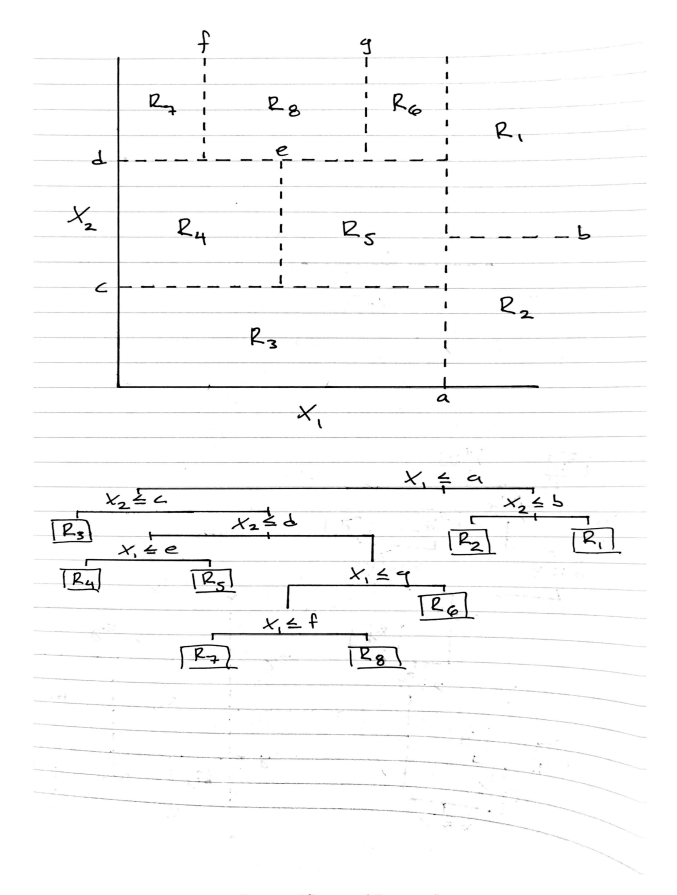
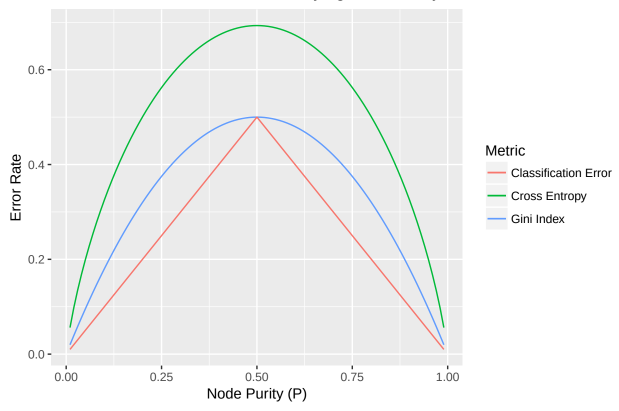


Figure 1: "Conceptual Exercise 1"  $\stackrel{}{2}$ 

```
suppressPackageStartupMessages(library(ggplot2))
pm1 \leftarrow seq(0.01, 0.99, 0.01)
# defining functions
two class gini <- function(prob) {</pre>
    return((prob * (1-prob)) + (prob * (1-prob)))
two_class_entropy <- function(prob) {</pre>
    return(-1*(prob*log(prob) + (1 - prob)*log(1 - prob)))
}
class_error <- function(prob) {</pre>
    return(1 - pmax(prob, (1 - prob)))
# error calculation
classification_error <- class_error(pm1)</pre>
gini_error <- two_class_gini(pm1)</pre>
entropy_error <- two_class_entropy(pm1)</pre>
# bring all data together for plotting
error_df <- data.frame(x = pm1,</pre>
                        gini = gini_error,
                        class = classification_error,
                        entropy = entropy_error)
# error plot
ggplot(error_df, aes(x = x, y = gini)) +
    geom_line(aes(color = 'Gini Index')) +
    geom_line(aes(y = entropy, color = 'Cross Entropy')) +
    geom_line(aes(y = class, color = 'Classification Error')) +
    guides(color = guide_legend(title = "Metric")) +
    ggtitle("Classification Error Rates with Varying Node Purity") +
    ylab("Error Rate") +
    xlab("Node Purity (P)")
```

## Classification Error Rates with Varying Node Purity



4

(sketch on following page)

5

```
probs <- c(0.1, 0.15, 0.2, 0.2, 0.55, 0.6, 0.6, 0.65, 0.7, 0.75)

# majority vote chooses red
majority_vote <- ifelse(probs >= 0.5, 'red', 'green')
table(majority_vote)

## majority_vote
## green red
## 4 6

# average chooses green
ifelse(mean(probs) >= 0.5, 'red', 'green')

## [1] "green"
```

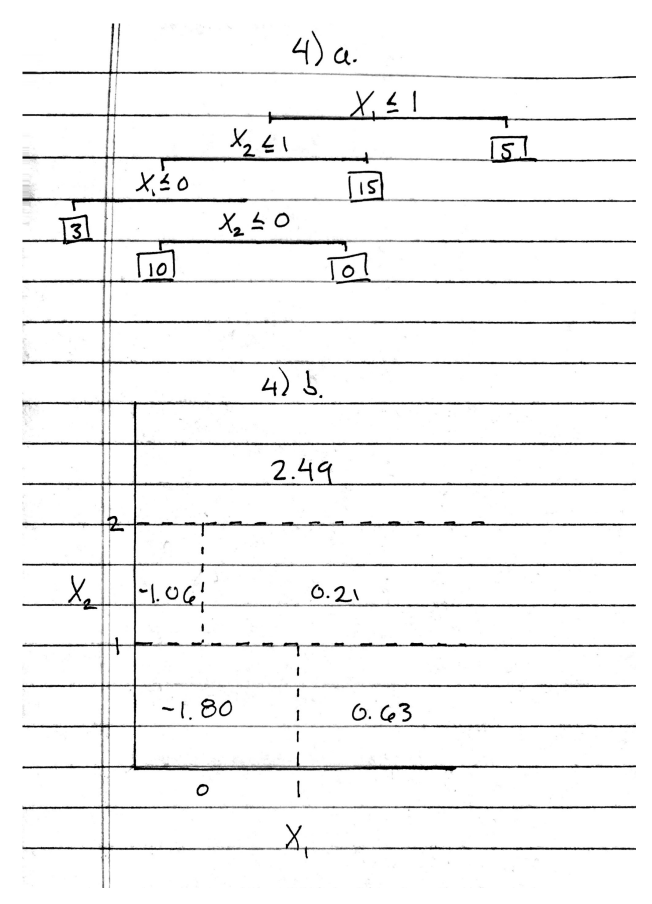


Figure 2: "Conceptual Exercise 4" 5

**Broad Strokes** - Use recursive binary splitting to continually split the data until there are fewer than x training observations in each terminal node, where x is predefined. The prediction for each terminal node will be the average of the response values of the training observations in that terminal node.

**Detail** - Starting with all the observations in the training set in one group, choose **the** value that, when split on, produces the lowest RSS among the two groups dictated by that split value.

- When the data set is split into two groups, the **single prediction** for the observations that are in group A will be the mean of the response values of the observations that are in group A, and the **single prediction** for the observations that are in group B will be the mean of the response values of the observations that are in group B.
- To calculate the (training) RSS of a particular split, sum the squared differences between the predicted value and the actual value of the response for all observations.

Now the data is parititioned into two groups. Repeat the same process again, treating each of those two groups as the parent group, which will be split into two child groups, leading to four total groups/divisions of the data set. Repeating again would lead to eight total division of the data set. Repeat this process until there are less than x training observations in each terminal node. Note that if only one of the parent groups has more than x training observations in it, but the other has less than x, only the group with more than x would be split further (in other words, the tree doesn't have to split evenly all the way down).

### Applied

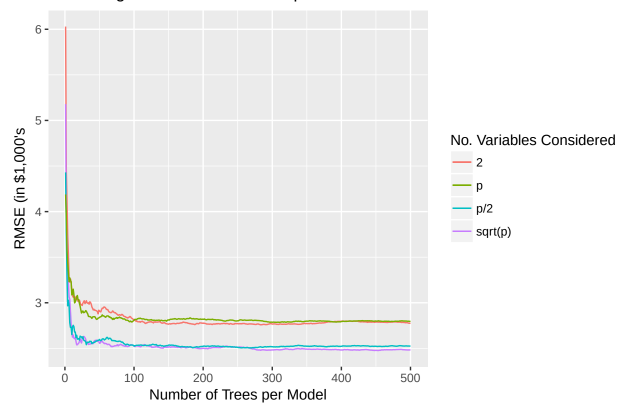
### 7

Looking at the plot below, it is clear that the testing error rate levels off after around 100 trees. Notably, as is characteristic of Random Forests, the classic "U" shape of the testing error is not present, showing a lack of overfitting as the number of trees in the ensemble increases.

```
# load libraries
suppressPackageStartupMessages(library(ISLR))
suppressPackageStartupMessages(library(MASS))
suppressPackageStartupMessages(library(randomForest))
attach(Boston)
set.seed(5)
# train test split
train <- sample(c(TRUE, FALSE), nrow(Boston), replace = TRUE)</pre>
X.train <- Boston[train, -14]</pre>
X.test <- Boston[-train, - 14]</pre>
y.train <- Boston[train, 14]</pre>
y.test <- Boston[-train, 14]</pre>
# modeling
rf.model.p <- randomForest(x = X.train,</pre>
                           y = y.train,
                           xtest = X.test,
                           ytest = y.test,
                           mtry = ncol(Boston) - 1,
```

```
ntree = 500)
rf.model.half_p <- randomForest(x = X.train,</pre>
                         y = y.train,
                         xtest = X.test,
                         ytest = y.test,
                         mtry = ceiling(ncol(Boston)/2),
                         ntree = 500)
rf.model.sqrt_p <- randomForest(x = X.train,</pre>
                         y = y.train,
                         xtest = X.test,
                         ytest = y.test,
                         mtry = ceiling(sqrt(ncol(Boston))),
                         ntree = 500)
rf.model.2 <- randomForest(x = X.train,</pre>
                         y = y.train,
                         xtest = X.test,
                         ytest = y.test,
                         mtry = 2,
                         ntree = 500)
# create dataframe
model.df <- data.frame(x = 1:500,
                       model_2 = rf.model.2$test$mse,
                       model_sqrt_p = rf.model.sqrt_p$test$mse,
                       model_half_p = rf.model.half_p$test$mse,
                       model_p = rf.model.p$test$mse)
# plot errors
ggplot(model.df, aes(x = x, y = model_2)) +
    geom_line(aes(y = sqrt(model_2), color = '2')) +
    geom_line(aes(y = sqrt(model_sqrt_p), color = 'sqrt(p)')) +
    geom_line(aes(y = sqrt(model_half_p), color = 'p/2')) +
    geom_line(aes(y = sqrt(model_p), color = 'p')) +
    guides(color = guide_legend(title = "No. Variables Considered")) +
    ggtitle("Predicting Median Owner-Occupied Homes") +
    xlab("Number of Trees per Model") +
    ylab("RMSE (in $1,000's")
```

## Predicting Median Owner-Occupied Homes



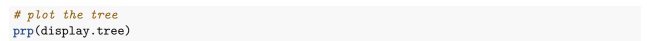
8

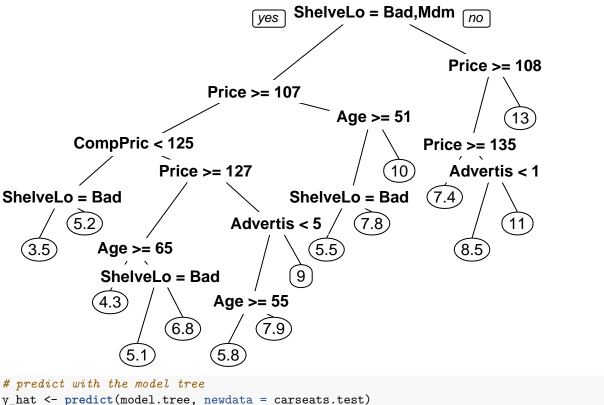
### • A.

```
detach(Boston)
attach(Carseats)

# train test split
set.seed(5)
train <- sample(c(TRUE, FALSE), nrow(Carseats), replace = TRUE)
carseats.train <- Carseats[train,]
carseats.test <- Carseats[-train,]</pre>
```

#### • B.





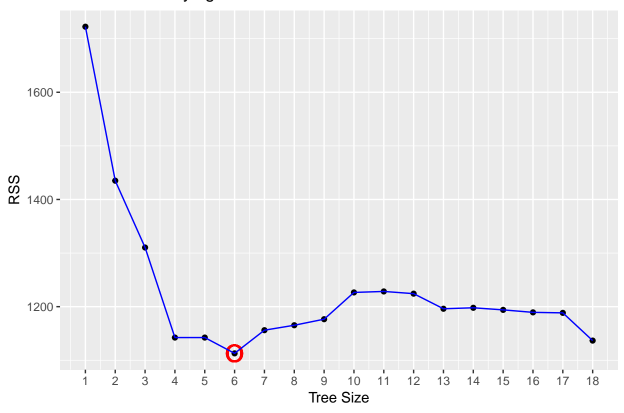
```
# predict with the model tree
y_hat <- predict(model.tree, newdata = carseats.test)
mse <- mean((carseats.test$Sales - y_hat)^2)
print(paste("Test MSE =", round(mse, 4)))</pre>
```

```
## [1] "Test MSE = 3.2113"
```

• C. Looking at the plot of the cross validated error against tree size below, it appears that pruning the tree would help, albiet minorly.

```
# cross validation
set.seed(2)
cv.carseats <- cv.tree(model.tree)</pre>
ggplot(data.frame(x = cv.carseats\$size,
                  y = cv.carseats$dev), aes(x, y)) +
    geom_point() +
    geom_point(x = cv.carseats$size[which.min(cv.carseats$dev)],
               y = cv.carseats$dev[which.min(cv.carseats$dev)],
               color = 'red',
               shape = '0',
               size = 6) +
    scale_x_continuous(breaks = rev(cv.carseats$size),
                       labels = rev(cv.carseats$size)) +
    geom_line(color = 'blue') +
    ggtitle("CV Error for Varying Tree Sizes") +
    xlab("Tree Size") +
    ylab("RSS")
```

### CV Error for Varying Tree Sizes

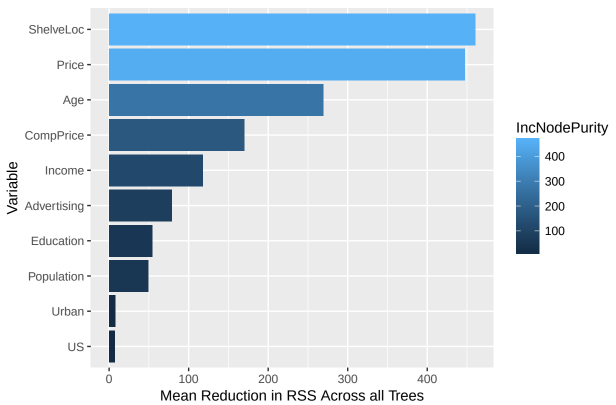


• **D**. The variable importance plot shown below indicates which variables were "most useful" in reducing child node impurity. In this case (regression), the *Increase in Node Purity* is calculated by finding the reduction in the RSS from splitting on that variable. For every split *in every tree* where x variable is split on, the reduction in the RSS is recorded, and then this is averaged over all trees.

```
# bagged decision trees
bag.model <- randomForest(Sales ~ .,</pre>
                           data = carseats.train,
                           mtry = ncol(carseats.train) - 1,
                           importance = TRUE)
y_hat <- predict(bag.model, newdata = carseats.test)</pre>
mse <- mean((carseats.test$Sales - y_hat)^2)</pre>
print(paste("Test MSE =", round(mse, 4)))
## [1] "Test MSE = 1.4539"
# variable importance plot
var imp df <- data.frame(importance(bag.model))</pre>
ggplot(var_imp_df, aes(x = reorder(row.names(var_imp_df),
                                     IncNodePurity),
                        y = IncNodePurity,
                        fill = IncNodePurity)) +
    geom_bar(stat = 'identity') +
    coord_flip() +
    ggtitle("Variable Importances for Bagged Decision Trees") +
    ylab("Mean Reduction in RSS Across all Trees") +
```

### xlab("Variable")

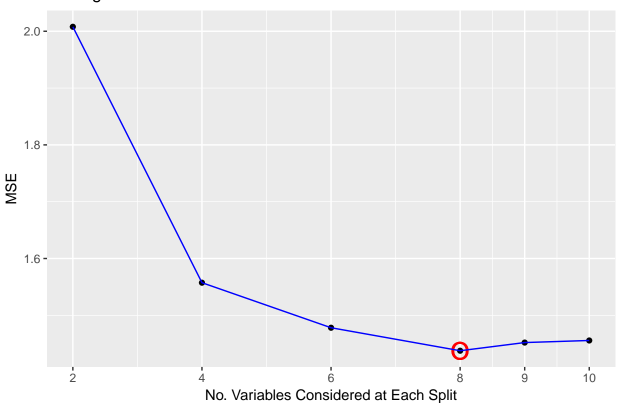
## Variable Importances for Bagged Decision Trees



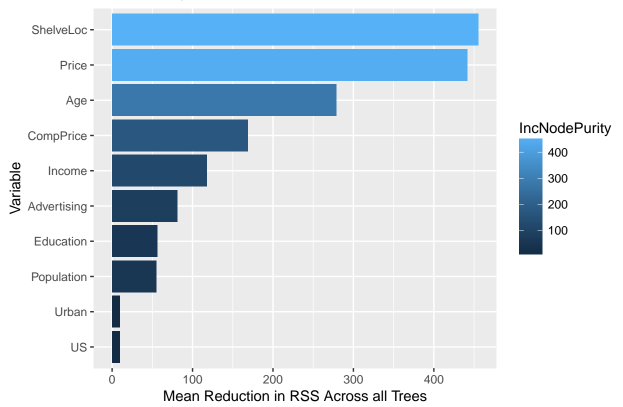
• E. Looking at the first plot below, it is clear that as the number of variables considered at each split increases, the testing MSE decreases up to a point (8, in this forest) and then increases again (although minorly).

```
# test various number of attributes considered at each split
set.seed(2)
m \leftarrow c(2, 4, 6, 8, 9, 10)
errors <- rep(0, length(m))
for (i in 1:length(m)) {
    rf.model <- randomForest(Sales ~ .,</pre>
                                data = carseats.train,
                                ntree = 500,
                                mtry = m[i],
                                importance = TRUE)
    y hat <- predict(rf.model, newdata = carseats.test)</pre>
    errors[i] <- mean((carseats.test$Sales - y_hat)^2)</pre>
}
# plot various errors
ggplot(data.frame(x = m,
                   y = errors), aes(x, y)) +
    geom_point() +
    geom_point(x = m[which.min(errors)],
```

## Testing MSE for Various M



## Variable Importances for Random Forest



9

#### • A.

```
detach(Carseats)
attach(OJ)

# train test split
set.seed(5)
train <- sample(1:dim(OJ)[1], 800)
oj.train <- OJ[train, ]
oj.test <- OJ[-train, ]</pre>
```

• B. The tree has 8 terminal nodes and has a misclassification error rate (given by  $\frac{\sum_{i=1}^{n}I(y_i\neq\hat{y_1})}{n}$ ) is 0.17 model.tree <- tree(Purchase ~ ., data = oj.train)

#### summary(model.tree)

```
##
## Classification tree:
## tree(formula = Purchase ~ ., data = oj.train)
## Variables actually used in tree construction:
## [1] "LoyalCH" "PriceDiff" "DiscCH"
## Number of terminal nodes: 7
## Residual mean deviance: 0.7786 = 617.4 / 793
## Misclassification error rate: 0.1838 = 147 / 800
```

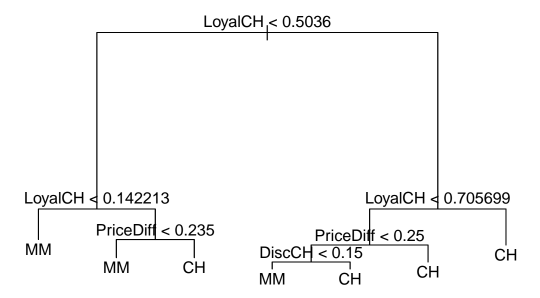
• C. Looking at node 4 (the first terminal node, denoted by an asterisk), it is shown that any observation with a *LoyalCH* value (numerical value symbolizing customer loyalty to the CH brand) less than 0.00589885 will be predicted to be MM (Minute Maid). There are 64 observations in the training data that are allocated to this node. There is a 94% probability of MM and conversely a 6% probability of CH.

#### model.tree

```
## node), split, n, deviance, yval, (yprob)
##
         * denotes terminal node
##
   1) root 800 1064.00 CH ( 0.61750 0.38250 )
##
##
      2) LoyalCH < 0.5036 354 435.50 MM ( 0.30508 0.69492 )
##
        4) LoyalCH < 0.142213 100
                                    45.39 MM ( 0.06000 0.94000 ) *
##
        5) LoyalCH > 0.142213 254
                                   342.20 MM ( 0.40157 0.59843 )
##
         10) PriceDiff < 0.235 136
                                   153.00 MM ( 0.25000 0.75000 ) *
         11) PriceDiff > 0.235 118 160.80 CH ( 0.57627 0.42373 ) *
##
##
      3) LoyalCH > 0.5036 446 352.30 CH ( 0.86547 0.13453 )
        6) LoyalCH < 0.705699 154 189.50 CH ( 0.69481 0.30519 )
##
         12) PriceDiff < 0.25 85 117.50 CH ( 0.52941 0.47059 )
##
##
           24) DiscCH < 0.15 77 106.60 MM ( 0.48052 0.51948 ) *
           25) DiscCH > 0.15 8
                                  0.00 CH ( 1.00000 0.00000 ) *
##
##
         13) PriceDiff > 0.25 69
                                   45.30 CH ( 0.89855 0.10145 ) *
        7) LoyalCH > 0.705699 292 106.30 CH ( 0.95548 0.04452 ) *
##
```

• **D**. As displayed by the first split on LoyalCH, the largest reduction in the misclassification rate is achieved by splitting on that variable. Further, both the second splits are LoyalCH to increase terminal node purity. For example, if an incoming observation has LoyalCh = 0.45 and PriceDiff = 0.5, that observation would "fall" down the left hand side of the top split, then right side of both the following splits.

```
plot(model.tree)
text(model.tree, pretty = 0)
```



• E. Looking at the confusion matrix below, the misclassification rate is 1 - Accuracy.

```
suppressPackageStartupMessages(library(caret))
y.hat <- predict(model.tree, newdata = oj.test, type = 'class')
cm <- confusionMatrix(y.hat, oj.test$Purchase)
error <- sum(cm$table[1,2], cm$table[2,1])/sum(cm$table)
print(paste("Misclassification Rate:", round(error, 4)))</pre>
```

## [1] "Misclassification Rate: 0.2444"

• F.

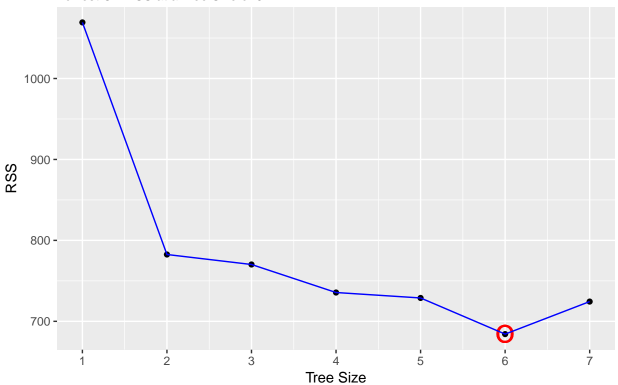
```
set.seed(5)
cv.oj <- cv.tree(model.tree)</pre>
```

#### • G & H.

```
ggplot(data.frame(x = cv.oj$size,
                  y = cv.oj\$dev), aes(x, y)) +
   geom_point() +
    geom point(x = cv.oj$size[which.min(cv.oj$dev)],
               y = cv.oj$dev[which.min(cv.oj$dev)],
               color = 'red',
               shape = '0',
               size = 6) +
    geom_line(color = 'blue') +
    scale_x_continuous(breaks = cv.oj$size,
                       labels = cv.oj$size) +
    ggtitle("CV Error for Varying Tree Sizes",
            paste("Lowest CV RSS at a Tree Size of",
                  cv.oj$size[which.min(cv.oj$dev)])) +
   xlab("Tree Size") +
   ylab("RSS")
```

### CV Error for Varying Tree Sizes

Lowest CV RSS at a Tree Size of 6



• I. (Although cross validation indicated a tree size of 6, I am setting the size to 4 in order to illustrate the point of the problem).

• J. The unpruned tree has a (slightly) lower training misclassification error rate than the pruned tree.

```
summary(pruned.tree)
```

## [1] "LoyalCH"

## Variables actually used in tree construction:

"PriceDiff" "DiscCH"

```
##
## Classification tree:
## snip.tree(tree = model.tree, nodes = 3L)
## Variables actually used in tree construction:
## [1] "LoyalCH" "PriceDiff"
## Number of terminal nodes: 4
## Residual mean deviance: 0.8938 = 711.4 / 796
## Misclassification error rate: 0.1875 = 150 / 800
summary(model.tree)
##
## Classification tree:
## tree(formula = Purchase ~ ., data = oj.train)
```

```
## Number of terminal nodes: 7
## Residual mean deviance: 0.7786 = 617.4 / 793
## Misclassification error rate: 0.1838 = 147 / 800
```

• **K**. As shown below, pruning the tree to a simpler pays off when it comes to testing data. The pruned tree has 78.52% accuracy while the fully grown tree has 75.56% accuracy.

```
y.hat.prune <- predict(pruned.tree, newdata = oj.test, type = 'class')
y.hat.normal <- predict(model.tree, newdata = oj.test, type = 'class')
cm.prune <- confusionMatrix(y.hat.prune, oj.test$Purchase)
cm.normal <- confusionMatrix(y.hat.normal, oj.test$Purchase)
cm.prune$overall[1]

## Accuracy
## 0.7851852
cm.normal$overall[1]</pre>
## Accuracy
## 0.7555556
```

10

• A.

```
detach(OJ)
attach(Hitters)

# transform data set
Hitters <- Hitters[complete.cases(Hitters$Salary),]
Hitters$Salary <- log(Hitters$Salary)</pre>
```

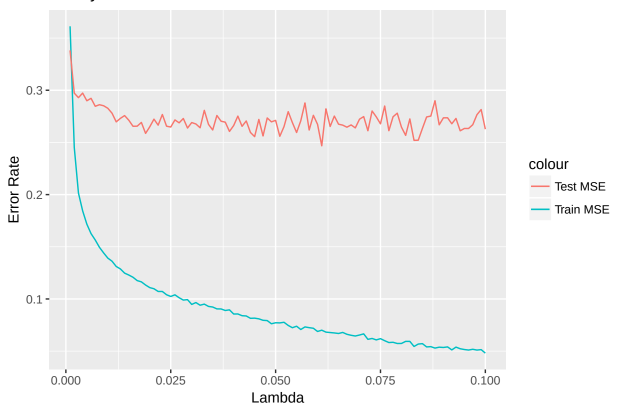
• B.

```
hitters.train <- Hitters[1:200,]
hitters.test <- Hitters[201:nrow(Hitters),]
```

• C & D.

```
train.y.hat <- predict(hitters.boost,</pre>
                      newdata = hitters.train,
                      n.trees = 1000)
    test.y.hat <- predict(hitters.boost,</pre>
                  newdata = hitters.test,
                  n.trees = 1000)
    train.errors[i] <- mean((train.y.hat - hitters.train$Salary)^2)</pre>
    test.errors[i] <- mean((test.y.hat - hitters.test$Salary)^2)</pre>
}
# plot results
ggplot(data.frame(x = lambdas,
                   train = train.errors,
                   test = test.errors), aes(x , test)) +
    geom_line(aes(y = train, color = "Train MSE")) +
    geom_line(aes(y = test, color = "Test MSE")) +
    ggtitle("Salary Prediction Errors") +
    xlab("Lambda") +
    ylab("Error Rate")
```

### Salary Prediction Errors



#### • E.

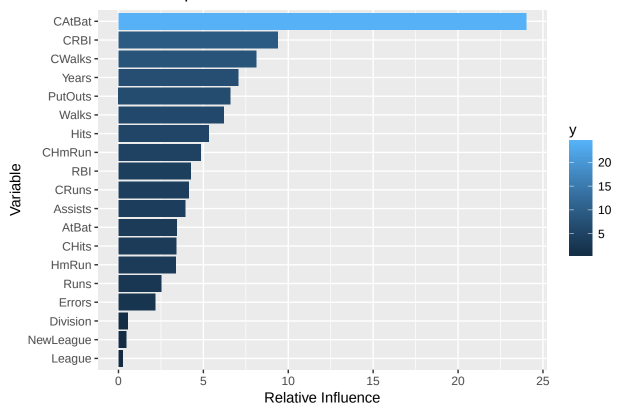
```
suppressPackageStartupMessages(library(glmnet))
# create training and testing matrices
x.train <- model.matrix(Salary ~ .,</pre>
```

```
data = hitters.train)[,-1]
y.train <- hitters.train$Salary</pre>
x.test <- model.matrix(Salary ~ .,</pre>
                         data = hitters.test)[,-1]
y.test <- hitters.test$Salary</pre>
# ridge regression
ridge.model <- glmnet(x.train,</pre>
                        y.train,
                       alpha = 0)
ridge.y.hat <- predict(ridge.model, newx = x.test)</pre>
ridge.mse <- mean((ridge.y.hat - y.test)^2)</pre>
# lasso regression
lasso.model <- glmnet(x.train,</pre>
                        y.train,
                        alpha = 1)
lasso.y.hat <- predict(lasso.model, newx = x.test)</pre>
lasso.mse <- mean((lasso.y.hat - y.test)^2)</pre>
# final boosted model
boosted.model <- gbm(Salary ~ .,</pre>
                      data = hitters.train,
                      distribution = 'gaussian',
                      n.trees = 1000,
                      shrinkage = lambdas[which.min(test.errors)])
boosted.y.hat <- predict(boosted.model,</pre>
                           newdata = hitters.test,
                          n.trees = 1000)
boosted.mse <- mean((boosted.y.hat - hitters.test$Salary)^2)</pre>
print(paste("Boosted regression trees testing MSE | ", round(boosted.mse, 4)))
## [1] "Boosted regression trees testing MSE | 0.2562"
print(paste("Lasso regression testing MSE | ", round(lasso.mse, 4)))
## [1] "Lasso regression testing MSE | 0.4756"
print(paste("Ridge regression testing MSE | ", round(ridge.mse, 4)))
## [1] "Ridge regression testing MSE | 0.5145"
```

• **F**. *CAtbat* is the most important variable in the boosted model, which equates to the number of times a player was at bat during his career.

### xlab("Variable")

## Variable Importances for Boosted Decision Trees



#### • G.

## [1] "Bagged regression trees testing MSE | 0.2352"

### 11

### • A.

```
detach(Hitters)
attach(Caravan)

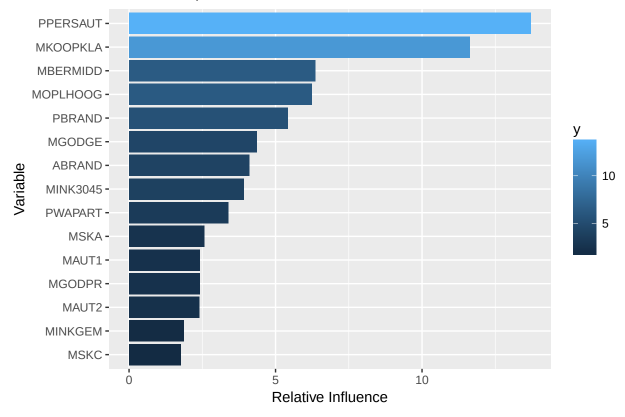
Caravan$Purchase <- ifelse(Caravan$Purchase == "Yes", 1, 0)

caravan.train <- Caravan[1:1000,]
caravan.test <- Caravan[1001:nrow(Caravan),]</pre>
```

#### • B.

```
set.seed(5)
# build boosted classification model
boost.caravan <- gbm(Purchase ~ . - PVRAAUT - AVRAAUT,
                     data = caravan.train,
                     distribution = 'gaussian',
                     n.trees = 1000,
                     shrinkage = 0.01)
# variable importance plot
ggplot(var_imp_df, aes(x = reorder(x, y),
                       y = y,
                       fill = y)) +
    geom_bar(stat = 'identity') +
    coord_flip() +
   ggtitle("Variable Importances for Boosted Decision Trees") +
   ylab("Relative Influence") +
   xlab("Variable")
```

### Variable Importances for Boosted Decision Trees



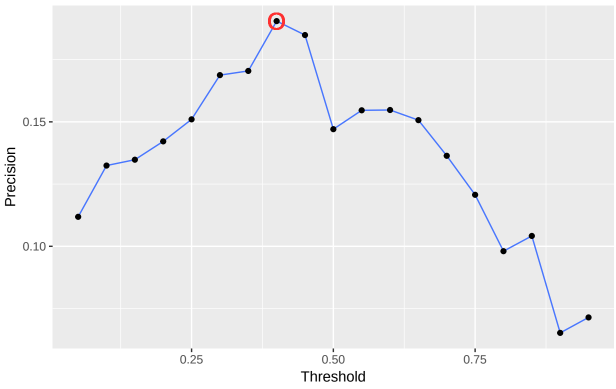
• C. While boosting produces a Positive Predictive Value (aka Precision) that is much higher than that of logistic regression, as shown below, one of the beauty's of probabilistic models is that has another conceptual knob to turn, toggling what the threshold is for a success. This is shown in the below plot, althought even the best precision outputted by logistic regression falls short of that of a boosted model

```
y.hat.prob <- predict(boost.caravan,</pre>
                      newdata = caravan.test,
                      n.trees = 1000,
                      type = 'response')
y.hat <- as.factor(ifelse(y.hat.prob > 0.2, 1, 0))
boost.summary <- confusionMatrix(as.factor(y.hat),</pre>
                      as.factor(caravan.test$Purchase),
                      positive = "1")
boost.summary$table
##
             Reference
## Prediction
                 0
            0 4494 277
##
               39
print(paste("Boosted ensemble precision = ", round(boost.summary$byClass[3], 4)))
## [1] "Boosted ensemble precision = 0.2353"
# Logistic Regression
log.mod <- glm(Purchase ~ .,</pre>
               data = caravan.train,
               family = 'binomial')
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
# prediction with various thresholds
log.mod.y.hat.logit <- predict(log.mod, newdata = caravan.test)</pre>
## Warning in predict.lm(object, newdata, se.fit, scale = 1, type =
## ifelse(type == : prediction from a rank-deficient fit may be misleading
log.mod.y.hat.prob <- exp(log.mod.y.hat.logit) / (exp(log.mod.y.hat.logit) + 1)</pre>
thresholds \leftarrow seq(0.05, 0.95, 0.05)
precisions <- rep(0, length(thresholds))</pre>
for (i in 1:length(thresholds)) {
    log.mod.y.hat <- ifelse(log.mod.y.hat.prob >= thresholds[i], 1, 0)
    log.mod.summary <- confusionMatrix(as.factor(log.mod.y.hat),</pre>
                                        as.factor(caravan.test$Purchase),
                                        positive = "1")
    precisions[i] <- log.mod.summary$byClass[3]</pre>
}
# plot precision
ggplot(data.frame(x = thresholds,
                  y = precisions), aes(x, y)) +
    geom_line(color = 'royalblue1') +
    geom_point() +
    geom_point(x = thresholds[which.max(precisions)],
               y = precisions[which.max(precisions)],
               shape = '0',
               size = 6,
               color = 'firebrick1') +
    ggtitle("Precision for Varying Thresholds",
            paste("Maximum Precision of", round(precisions[which.max(precisions)], 4), "at threshold of
```

```
xlab("Threshold") +
ylab("Precision")
```

## Precision for Varying Thresholds

Maximum Precision of 0.1905 at threshold of 0.4



### **12**

## alternate defaults.

Looking at the results below, of modeling whether the tomorrow's close price of GE will be higher than today's, it is shown that logistic regression outperforms the other models. Admittedly, with more robust features, I would imagine the scales would be tipped in favor of a more complex model, whether that be a Random Forest or Boosted Classification trees.

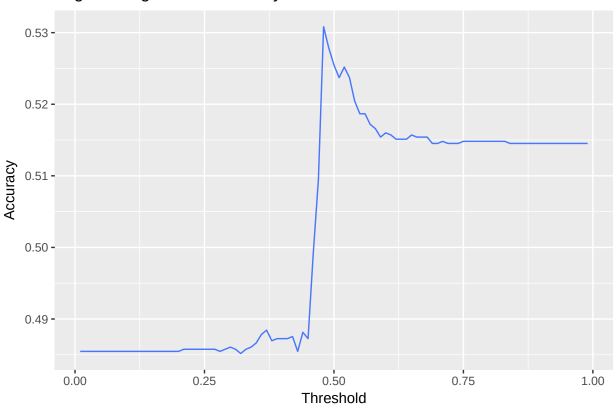
## 'loadSymbols' to automatically load data. getOption("getSymbols.env")
## and getOption("getSymbols.auto.assign") will still be checked for

```
##
## This message is shown once per session and may be disabled by setting
## options("getSymbols.warning4.0"=FALSE). See ?getSymbols for details.
## WARNING: There have been significant changes to Yahoo Finance data.
## Please see the Warning section of '?getSymbols.yahoo' for details.
## This message is shown once per session and may be disabled by setting
## options("getSymbols.yahoo.warning"=FALSE).
## [1] "GE"
# data cleaning
timestep_1 <- as.numeric(GE$GE.Close)</pre>
timestep_1 <- c(timestep_1, 0)</pre>
timestep_1 <- timestep_1[2:length(timestep_1)]</pre>
GE$timestep_plus_1 <- timestep_1</pre>
GE$Direction <- ifelse(GE$timestep_plus_1 > GE$GE.Close, 1, 0)
ge_df <- data.frame(GE)</pre>
ge_df <- ge_df %>% select(-one_of("timestep_plus_1"))
ge_df <- ge_df[1:nrow(ge_df) - 1,]</pre>
ge_df$Direction <- as.factor(ge_df$Direction)</pre>
# growth plot
chartSeries(GE)
```



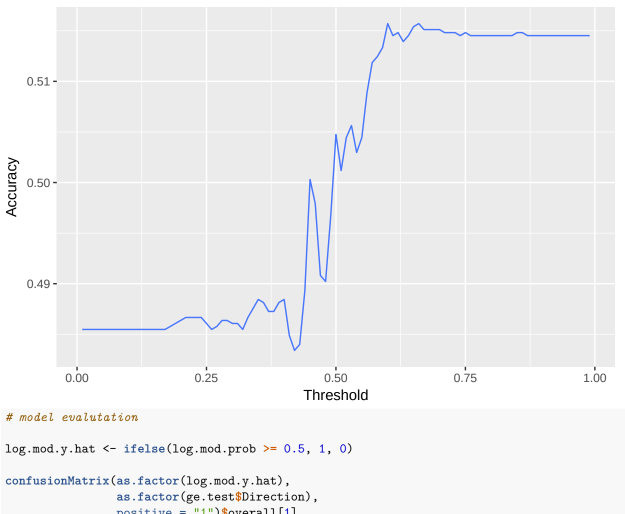
```
# train test splits
train.idx <- sample(c(TRUE, FALSE),</pre>
                      replace = TRUE,
                      nrow(ge_df),
                      prob = c(0.75, 0.25))
ge.train <- ge_df[train.idx, ]</pre>
ge.test <- ge_df[!train.idx, ]</pre>
x.train <- ge_df[train.idx, -7]</pre>
y.train <- ge_df[train.idx, 7]</pre>
x.test <- ge_df[!train.idx, -7]</pre>
y.test <- ge_df[!train.idx, 7]</pre>
# logistic regression
log.mod <- glm(Direction ~ .,</pre>
                data = ge.train,
                family = 'binomial')
log.mod.logit <- predict(log.mod, newdata = ge.test)</pre>
log.mod.prob <- exp(log.mod.logit) / (1 + exp(log.mod.logit))</pre>
# vary threholds
thresholds \leftarrow seq(0.01, 0.99, 0.01)
errors <- rep(0, length(thresholds))</pre>
for (i in 1:length(thresholds)) {
    log.mod.y.hat <- ifelse(log.mod.prob >= thresholds[i], 1, 0)
```

## Logistic Regression Accuracy for Various Thresholds



```
rf.mod.y.hat <- predict(rf.mod, newdata = ge.test)</pre>
# boosted trees
boost.mod <- gbm(as.character(Direction) ~ .,</pre>
                  data = ge.train,
                  distribution = 'bernoulli',
                  n.trees = 1000)
boost.mod.y.prob <- predict(boost.mod,</pre>
                            newdata = ge.test,
                            n.trees = 1000,
                            type = 'response')
boost_errors <- rep(0, length(thresholds))</pre>
# calculate accuracy for various thresholds of boosted model
for (i in 1:length(thresholds)) {
    boost.mod.y.hat <- ifelse(boost.mod.y.prob >= thresholds[i], 1, 0)
    suppressWarnings(
        cm <- confusionMatrix(as.factor(boost.mod.y.hat),</pre>
                               as.factor(ge.test$Direction),
                               positive = "1")
    )
    boost_errors[i] <- cm$overall[1]</pre>
# plot boosted model accuracy over various thresholds
ggplot(data.frame(x = thresholds,
                  y = boost_errors), aes(x, y)) +
    geom_line(color = 'royalblue1') +
    ggtitle("Boosted Accuracy for Various Thresholds") +
    ylab("Accuracy") +
    xlab("Threshold")
```

## **Boosted Accuracy for Various Thresholds**



```
log.mod.y.hat <- ifelse(log.mod.prob >= 0.5, 1, 0)
confusionMatrix(as.factor(log.mod.y.hat),
                positive = "1")$overall[1]
## Accuracy
## 0.525489
```

```
confusionMatrix(bagged.trees.y.hat,
                ge.test$Direction,
                positive = "1")$overall[1]
```

```
Accuracy
## 0.5047421
confusionMatrix(rf.mod.y.hat,
                ge.test$Direction,
```

positive = "1")\$overall[1]

```
## Accuracy
## 0.5106698
boost.mod.y.hat <- ifelse(boost.mod.y.prob >= 0.55, 1, 0)
confusionMatrix(as.factor(boost.mod.y.hat),
                as.factor(ge.test$Direction),
```

```
positive = "1")$overall[1]
```

## Accuracy ## 0.5044458