

Module 8: Solutions to Recommended Exercises

TMA4268 Statistical Learning V2021

Emma Skarstein, Michail Spitieris, Stefanie Muff
Department of Mathematical Sciences, NTNU

March 9, 2021

Problem 1 – Theoretical

a)

1. *Recursive binary splitting*: We find the best single partitioning of the data such that the reduction of RSS is the greatest. This process is applied sequentially to each of the split parts until a predefined minimum number of leave observation is reached.
2. *Cost complexity pruning* of the large tree from previous step, in order to obtain a sequence of best trees as a function of a parameter α . Each value of α corresponds to a subtree that minimize the following equation (several α s for the same tree):

$$\sum_{m=1}^{|T|} \sum_{i: x_i \in R_m} (y_i - \hat{y}_{R_m})^2 + \alpha |T|,$$

where $|T|$ is the number of terminal nodes.

3. *K-fold cross-validation* to choose α . For each fold:
 - Repeat Steps 1 and 2 on all but the k th folds of the training data.
 - Evaluate the mean squared prediction on the data in the left-out k th fold, as a function of α .
 - Average the results for each value of α and choose α to minimize the average error.
4. Return the subtree from Step 2 that corresponds to the chosen value of α .

For a **classification** tree we replace RSS with Gini index or entropy.

b)

Advantages

- Very easy to explain
- Can be displayed graphically
- Can handle both quantitative and qualitative predictors without the need to create dummy variables

Disadvantages

- The predictive accuracy is usually not very high
- They are non-robust. That is a small change in the data can cause a large change in the estimated tree

c)

Decision trees suffer from high variance. Recall that if we have B *i.i.d* observations of a random variable X with the same mean and variance σ^2 . We calculate the mean $\bar{X} = \frac{1}{B} \sum_{b=1}^B X_b$, and the variance of the mean is $\text{Var}(\bar{X}) = \frac{\sigma^2}{B}$. That is by averaging we get reduced variance.

For decision trees, if we have B training sets, we could estimate $\hat{f}_1(\mathbf{x}), \hat{f}_2(\mathbf{x}), \dots, \hat{f}_B(\mathbf{x})$ and average them as

$$\hat{f}_{avg}(\mathbf{x}) = \frac{1}{B} \sum_{b=1}^B \hat{f}_b(\mathbf{x}) .$$

However we do not have many data independent data sets, and we bootstrap to create B datasets. These datasets are however not completely independent and the reduction in variance is therefore not as large as for independent training sets.

To make the different trees that are built from each bootstrapped dataset more different, random forests use a random subset of the predictors to split the tree into new branches at each step. This decorrelates the different trees that are built from the B bootstrapped datasets, and consequently reduces variance.

d) An OOB is the set of observations that were not chosen to be in a specific bootstrap sample. From RecEx5-Problem 4c we have that on average $1 - 0.632 = 0.368$ are included in the OOB sample.

e)

Variable importance based on node impurity

Regression Trees: The total amount that the RSS is decreased due to splits of each predictor, averaged over the B trees.

Classification Trees: The importance is the mean decrease (over all B trees) in the Gini index by splits of a predictor.

Variable importance based on randomization

This measure is based on how much the predictive accuracy (MSE or gini index) is decreased when the variable is replaced by a permuted version of it. You find a drawing [here](#).

Problem 2 – Regression (Book Ex. 8)

a)

```
library(ISLR)
data("Carseats")
set.seed(4268)
n = nrow(Carseats)
train = sample(1:n, 0.7 * nrow(Carseats), replace = F)
test = (1:n)[-train]
Carseats.train = Carseats[train, ]
Carseats.test = Carseats[-train, ]
```

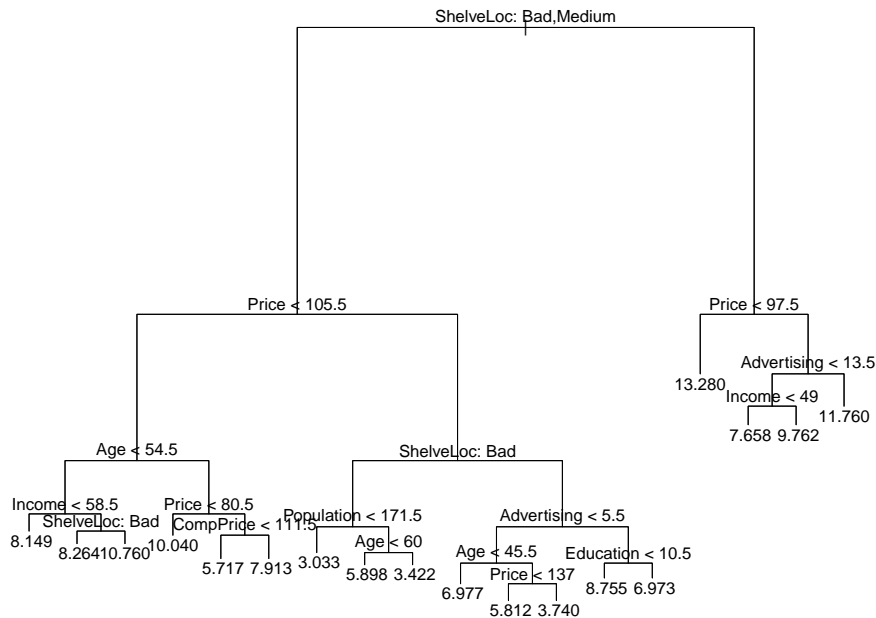
b)

```
library(tree)
tree.mod = tree(Sales ~ ., Carseats, subset = train)
summary(tree.mod)
```

```
##
## Regression tree:
## tree(formula = Sales ~ ., data = Carseats, subset = train)
## Variables actually used in tree construction:
## [1] "ShelveLoc" "Price" "Age" "Income" "CompPrice"
```

```
## [6] "Population" "Advertising" "Education"
## Number of terminal nodes: 18
## Residual mean deviance: 2.609 = 683.6 / 262
## Distribution of residuals:
##      Min. 1st Qu.  Median    Mean 3rd Qu.    Max.
## -3.74000 -1.12400 -0.06522  0.00000  1.06800  4.47200
```

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

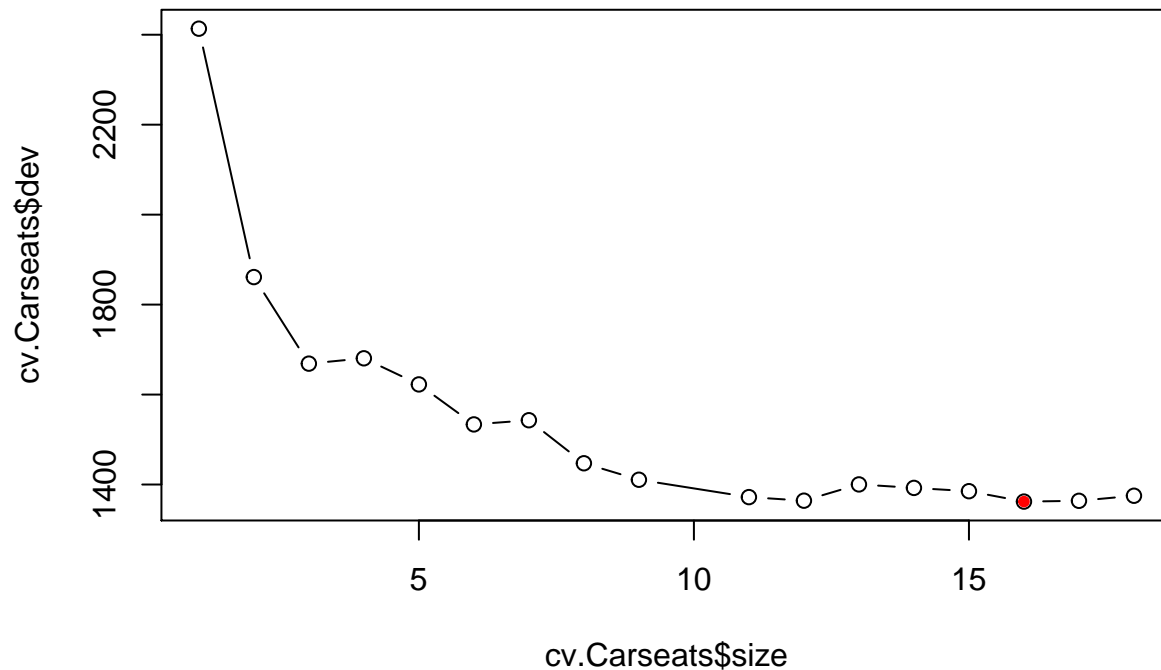


```
yhat = predict(tree.mod, newdata = Carseats.test)
mse = mean((yhat - Carseats.test$Sales)^2)
mse
```

```
## [1] 4.585249
```

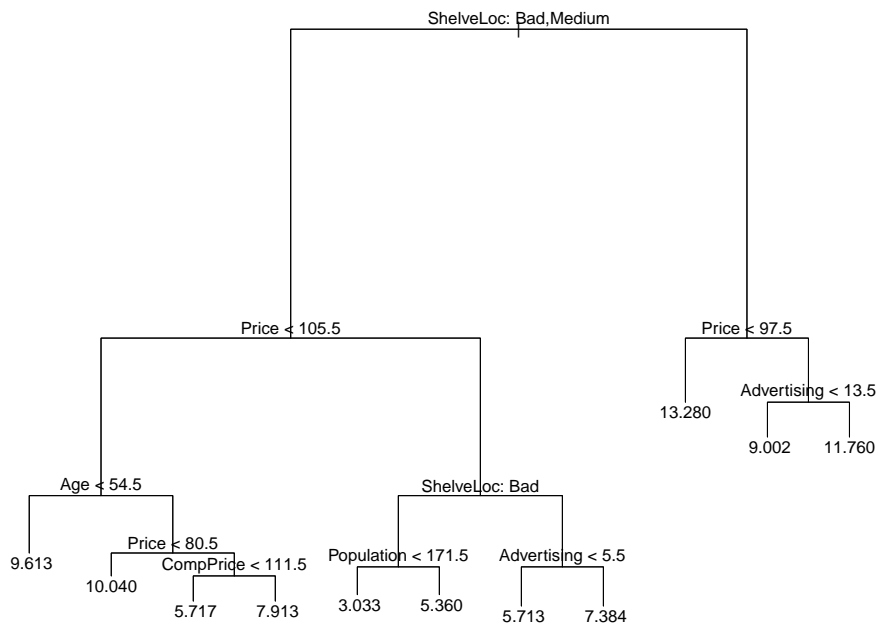
c)

```
set.seed(4268)
cv.Carseats = cv.tree(tree.mod)
tree.min = which.min(cv.Carseats$dev)
best = cv.Carseats$size[tree.min]
plot(cv.Carseats$size, cv.Carseats$dev, type = "b")
points(cv.Carseats$size[tree.min], cv.Carseats$dev[tree.min], col = "red", pch = 20)
```



We see that trees with sizes 11, 12, 16 and 17 have similar deviance values. We should ideally choose the tree of size 11 as it gives the simpler tree.

```
pr.tree = prune.tree(tree.mod, best = 11)
plot(pr.tree)
text(pr.tree, pretty = 0)
```



```
yhat = predict(pr.tree, newdata = Carseats.test)
mse = mean((yhat - Carseats.test$Sales)^2)
mse
```

```
## [1] 4.378499
```

There is a slight reduction in MSE for the pruned tree with 11 leaves.

d)

```
library(randomForest)
dim(Carseats)
```

```
## [1] 400 11
```

```
bag.Carseats = randomForest(Sales ~ ., Carseats.train, mtry = ncol(Carseats) - 1,
                             ntree = 500, importance = TRUE)
yhat.bag = predict(bag.Carseats, newdata = Carseats.test)
mse.bag = mean((yhat.bag - Carseats.test$Sales)^2)
mse.bag
```

```
## [1] 2.122958
```

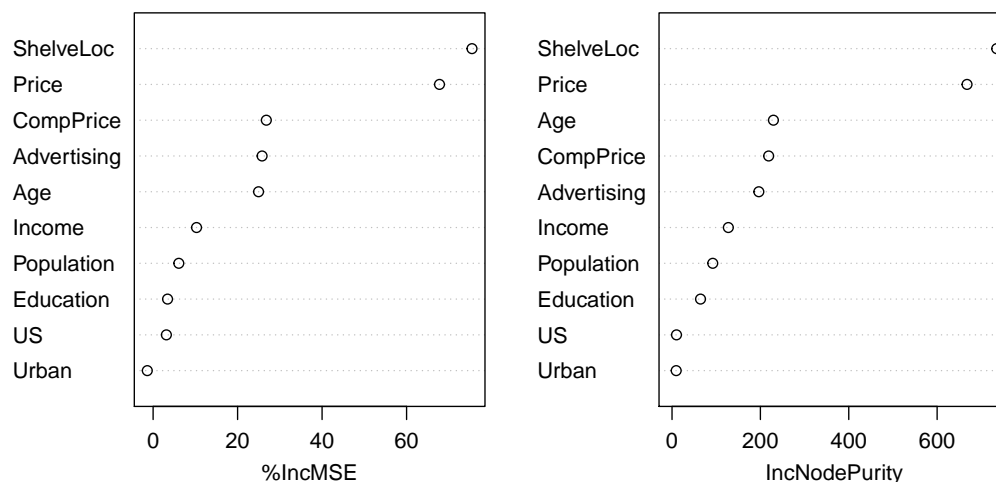
Bagging decreases the test MSE significantly to 2.12. From the importance plots we might conclude that Price and ShelfLoc are the most important Variables.

```
importance(bag.Carseats)
```

	%IncMSE	IncNodePurity
## CompPrice	26.803869	218.740455
## Income	10.284817	127.447480
## Advertising	25.795425	196.438893
## Population	6.084270	92.149065
## Price	67.791459	667.696518
## ShelfLoc	75.485534	734.902022
## Age	24.961130	229.491494
## Education	3.423565	64.510742
## Urban	-1.373635	9.423406
## US	3.141449	10.105870

```
varImpPlot(bag.Carseats)
```

bag.Carseats



e)

```
rf.Carseats = randomForest(Sales ~ ., data = Carseats.train, mtry = 3, ntree = 500,
                             importance = TRUE)
```

```
yhat.rf = predict(rf.Carseats, newdata = Carseats.test)
mse_forest <- mean((yhat.rf - Carseats.test$Sales)^2)
mse_forest
```

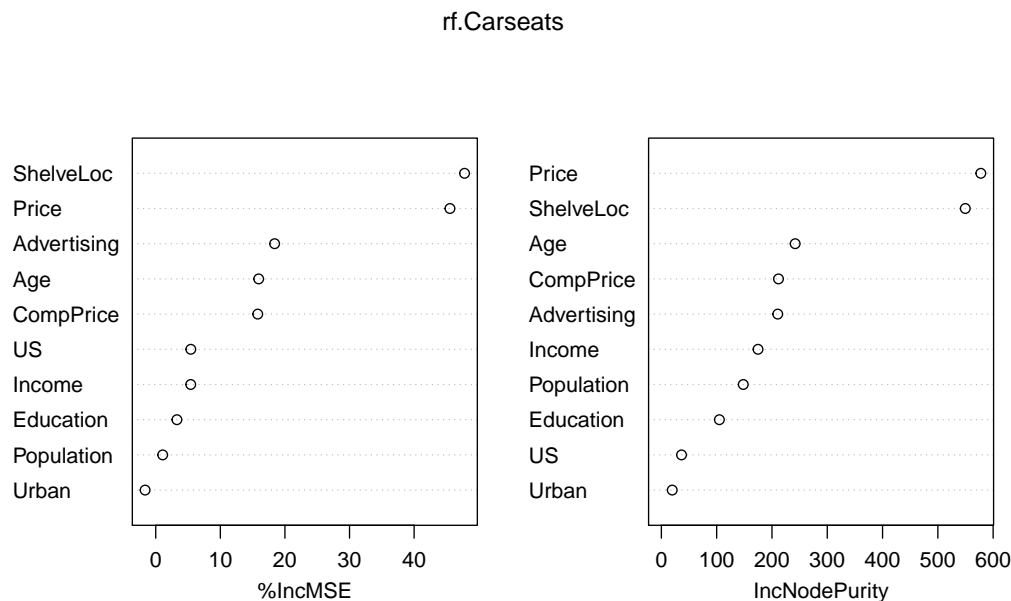
```
## [1] 2.25397
```

We use $\sqrt{p} = \sqrt{10} \approx 3$ trees, and we obtain an MSE of 2.25 which is slightly larger than Bagging MSE. The two most important Variables are again Price and ShelfLoc.

```
importance(rf.Carseats)
```

```
##           %IncMSE IncNodePurity
## CompPrice 15.789484    211.79213
## Income    5.415374    174.79625
## Advertising 18.402600    210.47149
## Population 1.076874    148.09993
## Price     45.548596    577.68865
## ShelfLoc  47.810006    549.62278
## Age       15.936114    241.99130
## Education  3.275725    104.89503
## Urban     -1.646580     19.63668
## US        5.427599     36.45647
```

```
varImpPlot(rf.Carseats)
```



f)

```
library(gbm)
r.boost = gbm(Sales ~ ., Carseats.train, distribution = "gaussian", n.trees = 500,
              interaction.depth = 4, shrinkage = 0.1)
yhat.boost = predict(r.boost, newdata = Carseats.test, n.trees = 500)
mse_boost <- mean((yhat.boost - Carseats.test$Sales)^2)
mse_boost
```

```
## [1] 2.151292
```

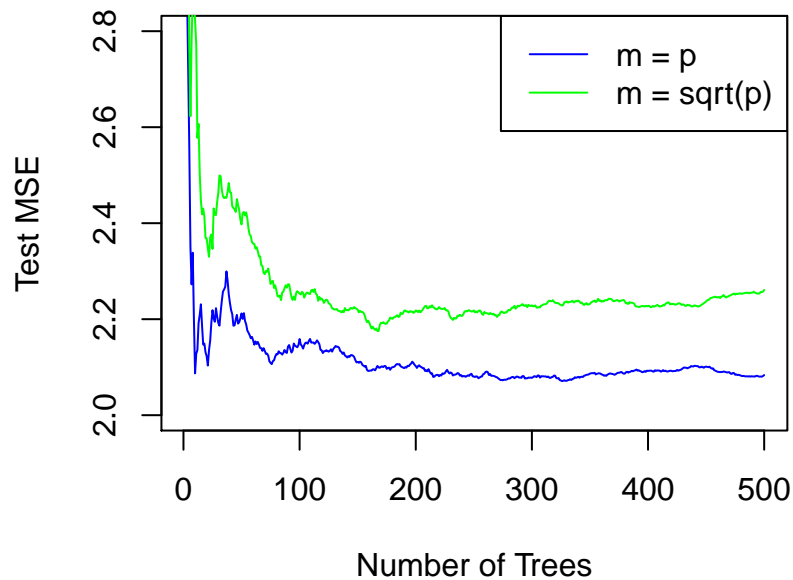
We see a further decrease in MSE by boosting our trees.

g)

```
train.predictors = Carseats.train[, -1]
test.predictors = Carseats.test[, -1]
Y.train = Carseats.train[, 1]
Y.test = Carseats.test[, 1]

bag.Car = randomForest(train.predictors, y = Y.train, xtest = test.predictors, ytest = Y.test,
  mtry = 10, ntree = 500)
rf.Car = randomForest(train.predictors, y = Y.train, xtest = test.predictors, ytest = Y.test,
  mtry = 3, ntree = 500)
plot(1:500, bag.Car$test$mse, col = "blue", type = "l", xlab = "Number of Trees",
  ylab = "Test MSE", ylim = c(2, 2.8))
lines(1:500, rf.Car$test$mse, col = "green")

legend("topright", c("m = p", "m = sqrt(p)"), col = c("blue", "green"), cex = 1,
  lty = 1)
```



Problem 3 – Classification

```
library(kernlab)
data(spam)
```

a)

Do this in R and read the Description.

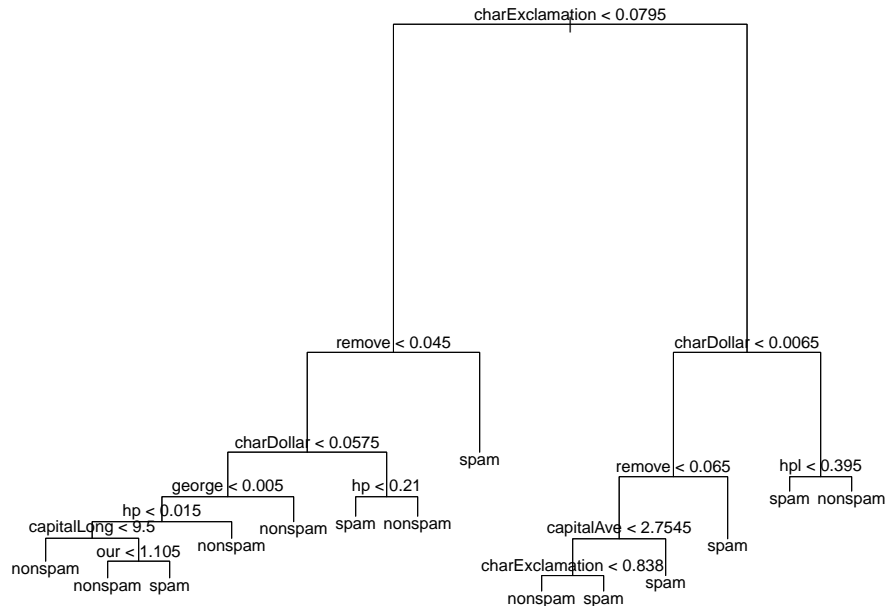
b)

```
library(ISLR)
set.seed(4268)
n = nrow(spam)
train = sample(1:n, 0.7 * n, replace = F)
test = (1:n)[-train]
spam.train = spam[train, ]
spam.test = spam[-train, ]
```

c)

```
spam.tree = tree(type ~ ., spam, subset = train)
```

```
plot(spam.tree)
text(spam.tree, pretty = 1)
```



```
summary(spam.tree)
```

```
##
## Classification tree:
## tree(formula = type ~ ., data = spam, subset = train)
## Variables actually used in tree construction:
## [1] "charExclamation" "remove" "charDollar" "george"
## [5] "hp" "capitalLong" "our" "capitalAve"
## [9] "hpl"
## Number of terminal nodes: 14
## Residual mean deviance: 0.4801 = 1539 / 3206
## Misclassification error rate: 0.08975 = 289 / 3220
```

d)

```
yhat = predict(spam.tree, spam[test, ], type = "class")
response.test = spam$type[test]
```

```
misclass = table(yhat, response.test)
misclass
```

```
##           response.test
## yhat      nonspam spam
## nonspam    781   67
## spam       67  466
```

```
1 - sum(diag(misclass))/sum(misclass)
```

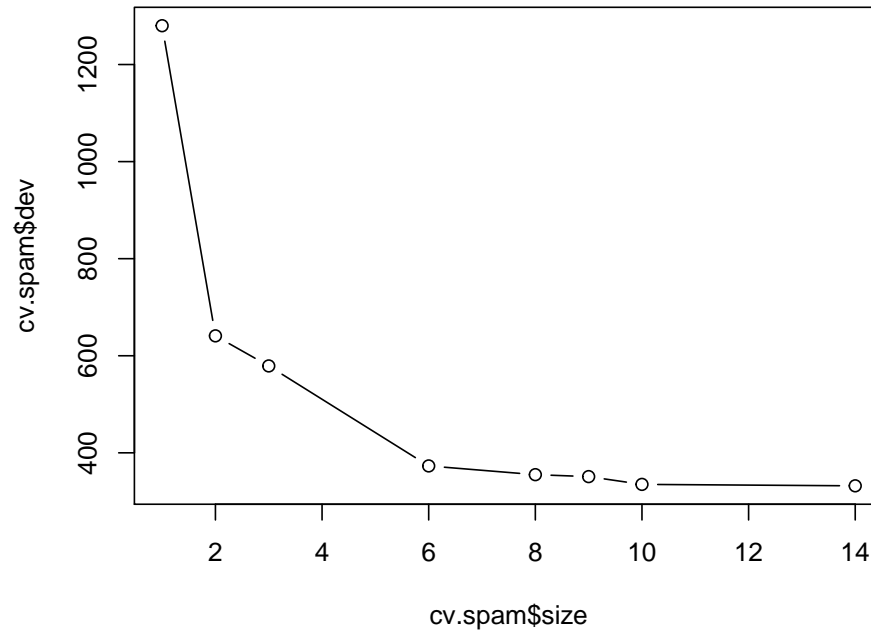
```
## [1] 0.09703114
```

e)


```
set.seed(4268)

cv.spam = cv.tree(spam.tree, FUN = prune.misclass)

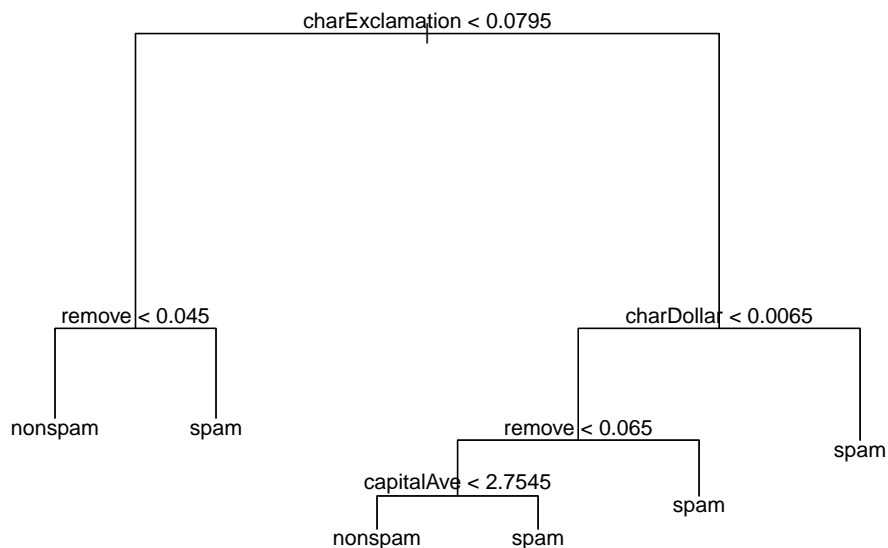
plot(cv.spam$size, cv.spam$dev, type = "b")
```



According to the plot the optimal number of terminal nodes is 6 (or larger). We choose 6 as this gives the simplest tree, and prune the tree according to this value.

```
prune.spam = prune.misclass(spam.tree, best = 6)

plot(prune.spam)
text(prune.spam, pretty = 1)
```



We predict the response for the test data:

```
yhat.prune = predict(prune.spam, spam[test, ], type = "class")
```

```
misclass.prune = table(yhat.prune, response.test)
misclass.prune
```

```
##           response.test
## yhat.prune nonspam spam
##   nonspam      796  104
##    spam        52  429
```

The misclassification rate is

```
1 - sum(diag(misclass.prune))/sum(misclass.prune)
```

```
## [1] 0.1129616
```

f)

```
library(randomForest)
bag.spam = randomForest(type ~ ., data = spam, subset = train, mtry = ncol(spam) -
  1, ntree = 500, importance = TRUE)
```

We predict the response for the test data as before:

```
yhat.bag = predict(bag.spam, newdata = spam[test, ])
```

```
misclass.bag = table(yhat.bag, response.test)
misclass.bag
```

```
##           response.test
## yhat.bag  nonspam spam
##   nonspam      810  43
##    spam        38  490
```

The misclassification rate is

```
1 - sum(diag(misclass.bag))/sum(misclass.bag)
```

```
## [1] 0.05865315
```

g)

We now use the random forest-algorithm and consider only $\sqrt{57} \approx 8$ of the predictors at each split. This is specified in mtry.

```
set.seed(4268)
rf.spam = randomForest(type ~ ., data = spam, subset = train, mtry = round(sqrt(ncol(spam) -
  1)), ntree = 500, importance = TRUE)
```

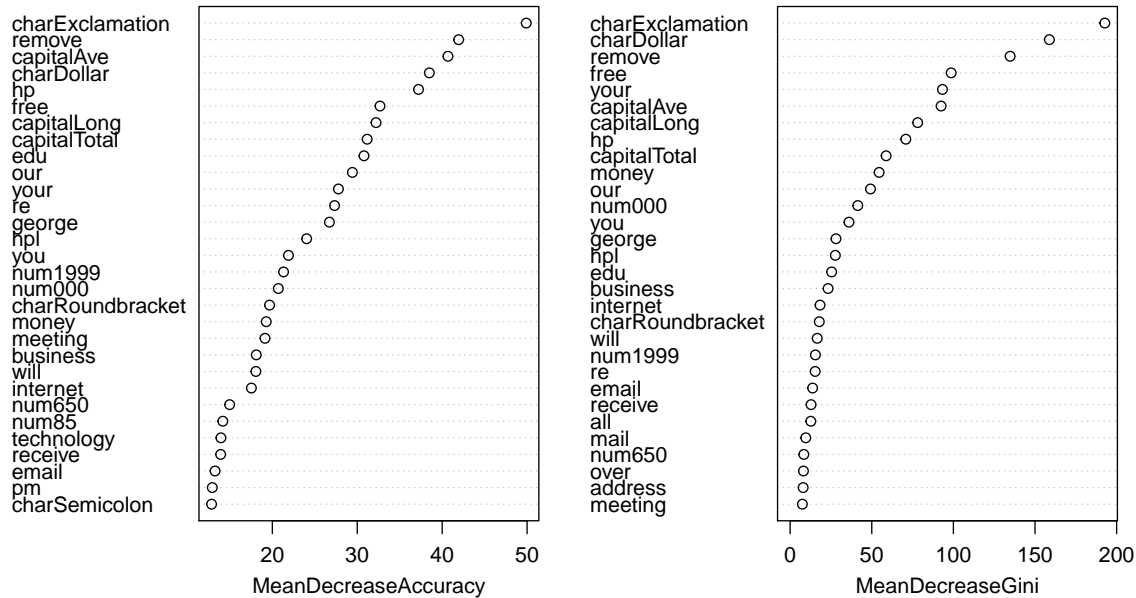
We study the importance of each variable

```
importance(rf.spam)
```

If MeanDecreaseAccuracy and MeanDecreaseGini are large, the corresponding covariate is important.

```
varImpPlot(rf.spam)
```

rf.spam



In this plot we see that **charExclamation** is the most important covariate, followed by **remove** and **charDollar**. This is as expected as these variables are used in the top splits in the classification trees we have seen so far.

We now predict the response for the test data.

```
yhat.rf = predict(rf.spam, newdata = spam[test, ])
```

```
misclass.rf = table(yhat.rf, response.test)
1 - sum(diag(misclass.rf))/sum(misclass.rf)
```

```
## [1] 0.044895
```

The misclassification rate is given by

```
misclass.rf
```

```
##           response.test
## yhat.rf  nonspam spam
## nonspam    824   38
## spam       24  495
```

h)

The `gbm()` function does not allow factors, so we have to use '1' and '0' instead of `spam` and `nonspam`

```
library(gbm)
set.seed(4268)

spamboost = spam
spamboost$type = c()
spamboost$type[spam$type == "spam"] = 1
spamboost$type[spam$type == "nonspam"] = 0

boost.spam = gbm(type ~ ., data = spamboost[train, ], distribution = "bernoulli",
  n.trees = 5000, interaction.depth = 3, shrinkage = 0.001)
```

We predict the response for the test data

```
yhat.boost = predict(boost.spam, newdata = spamboost[-train, ], n.trees = 5000, distribution = "bernoulli",  
  type = "response")
```

```
yhat.boost = ifelse(yhat.boost > 0.5, 1, 0) #Transform to 0 and 1 (nonspam and spam).
```

```
misclass.boost = table(yhat.boost, spamboost$type[test])
```

```
misclass.boost
```

```
##
```

```
## yhat.boost    0    1
```

```
##           0 812  52
```

```
##           1  36 481
```

and the misclassification rate is

```
1 - sum(diag(misclass.boost))/sum(misclass.boost)
```

```
## [1] 0.06372194
```

i)

We get lower missclassification rates for bagging, random forest and boosting than for a simple tree, which is expected.