# Titanic Survivor Prediction

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## Titanic Survivor Prediction - Classification

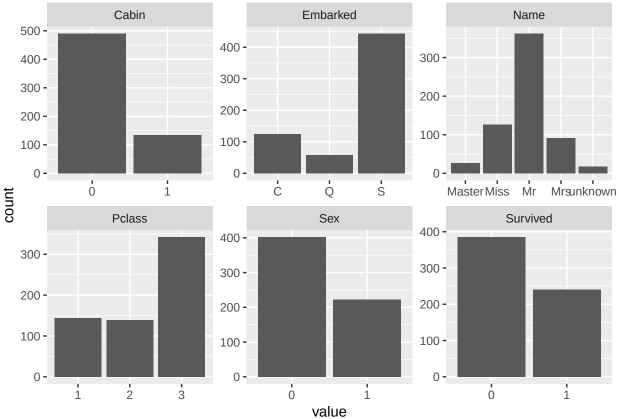
### Importing and Manipulating Data - Feature Engineering

```
library(caret)
## Loading required package: lattice
## Loading required package: ggplot2
## Registered S3 methods overwritten by 'ggplot2':
##
    method
                    from
##
     [.quosures
                    rlang
##
     c.quosures
                    rlang
    print.quosures rlang
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(MASS)
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
##
       select
#train and test
train <- read.csv("Datasets/train.csv", stringsAsFactors = TRUE, na.strings = "")</pre>
test <- read.csv("Datasets/test.csv", stringsAsFactors = TRUE)</pre>
#creating survived variables in test set
test$Survived <- NA
dat <- rbind(train,test)</pre>
summary(dat)
    PassengerId
                      Survived
                                        Pclass
                 Min. :0.0000
                                    Min. :1.000
## Min. : 1
## 1st Qu.: 328
                 1st Qu.:0.0000
                                    1st Qu.:2.000
```

```
Median: 655
                  Median :0.0000
                                   Median :3.000
##
   Mean : 655
                 Mean :0.3838
                                   Mean
                                          :2.295
   3rd Qu.: 982
                  3rd Qu.:1.0000
                                   3rd Qu.:3.000
## Max. :1309
                          :1.0000
                                   Max. :3.000
                  Max.
##
                  NA's
                          :418
##
                                 Name
                                               Sex
                                                              Age
## Connolly, Miss. Kate
                                       2
                                          female:466
                                                         Min. : 0.17
                                   :
## Kelly, Mr. James
                                       2
                                                         1st Qu.:21.00
                                    :
                                           male :843
## Abbing, Mr. Anthony
                                       1
                                                         Median :28.00
## Abbott, Mr. Rossmore Edward
                                       1
                                                         Mean :29.88
## Abbott, Mrs. Stanton (Rosa Hunt):
                                       1
                                                         3rd Qu.:39.00
## Abelson, Mr. Samuel
                                       1
                                                         Max.
                                                                :80.00
                                    :1301
## (Other)
                                                         NA's
                                                                :263
##
       SibSp
                        Parch
                                         Ticket
                                                         Fare
## Min. :0.0000
                    Min. :0.000
                                     CA. 2343: 11
                                                    Min. : 0.000
                                                    1st Qu.: 7.896
##
   1st Qu.:0.0000
                    1st Qu.:0.000
                                     1601
                                                8
## Median :0.0000
                    Median :0.000
                                     CA 2144 :
                                                    Median: 14.454
                                                 8
## Mean :0.4989
                    Mean :0.385
                                     3101295 :
                                                7 Mean : 33.295
## 3rd Qu.:1.0000
                    3rd Qu.:0.000
                                     347077 :
                                                    3rd Qu.: 31.275
## Max. :8.0000
                    Max. :9.000
                                     347082 : 7
                                                    Max. :512.329
##
                                     (Other) :1261
                                                    NA's
                                                          :1
##
                Cabin
                         Embarked
                         C :270
##
                   :327
                   : 6
## C23 C25 C27
                         Q
                              :123
## B57 B59 B63 B66: 5
                              :914
                         S
## G6
                   : 5
                         NA's: 2
## B96 B98
                   : 4
   (Other)
                   :275
## NA's
                   :687
#convert survived and pclass to factor variable
dat$Survived <- as.factor(dat$Survived)</pre>
dat$Pclass <- as.factor(dat$Pclass)</pre>
#Cabin Na values -> 0, otherwise 1
dat$Cabin <- as.factor(ifelse(is.na(dat$Cabin), 0, 1))</pre>
\#Gender \rightarrow male = 0, female = 1
dat$Sex <- as.factor(ifelse(dat$Sex == "male", 0, 1))</pre>
#NA values -> mean
dat$Age[is.na(dat$Age)] <- mean(dat$Age, na.rm=TRUE)</pre>
dat$Fare[is.na(dat$Fare)] <- mean(dat$Fare, na.rm=TRUE)</pre>
dat$Embarked[is.na(dat$Embarked)] <- "S"</pre>
#family size (if family = 1, then it's alone)
dat$family <- dat$SibSp + dat$Parch + 1</pre>
#converting names
name <- dat$Name
name <- sub(".*, ", "", name)
name <- sub("\\..*", "", name)
```

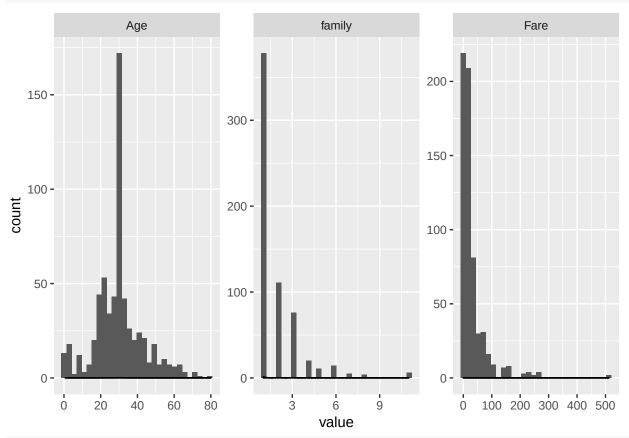
```
head(name)
## [1] "Mr"
              "Mrs" "Miss" "Mrs"
                                           "Mr"
                                   "Mr"
dat$Name <- as.character(name)</pre>
dat$Name (dat$Name %in% c("Capt", "Col", "Don", "Dr", "Jonkheer", "Lady", "Major", "Mile", "Mme", "Ms", "Rev", "S
dat$Name <- as.factor(dat$Name)</pre>
table(dat$Name)
##
## Master
                               Mrs unknown
              Miss
                        \mathtt{Mr}
               260
                       757
                               197
##
       61
#dropping variables
dat <- subset(dat, select = -c(PassengerId, SibSp, Parch, Ticket))</pre>
names (dat)
## [1] "Survived" "Pclass"
                              "Name"
                                         "Sex"
                                                    "Age"
                                                                "Fare"
## [7] "Cabin"
                  "Embarked" "family"
summary(dat)
## Survived
              Pclass
                            Name
                                     Sex
                                                   Age
## 0
       :549
              1:323 Master: 61
                                     0:843
                                             Min. : 0.17
               2:277 Miss
                                              1st Qu.:22.00
## 1
        :342
                              :260
                                      1:466
## NA's:418 3:709
                       \mathtt{Mr}
                              :757
                                              Median :29.88
##
                       Mrs
                              :197
                                              Mean
                                                     :29.88
##
                       unknown: 34
                                              3rd Qu.:35.00
##
                                              Max.
                                                     :80.00
##
         Fare
                      Cabin
                              Embarked
                                            family
## Min. : 0.000
                                              : 1.000
                      0:687
                              C:270 Min.
## 1st Qu.: 7.896
                      1:622
                              Q:123
                                        1st Qu.: 1.000
## Median : 14.454
                              S:916
                                        Median : 1.000
## Mean : 33.295
                                        Mean : 1.884
                                        3rd Qu.: 2.000
## 3rd Qu.: 31.275
## Max.
          :512.329
                                        Max.
                                               :11.000
train1 <- dat[1:891,] #train.csv</pre>
test1 <- dat[892:1309,] #test.csv
set.seed(125)
#cv splits
split<-createDataPartition(y=train1$Survived,p=0.7,list=FALSE)</pre>
training <- train1[split,] #training set of train data</pre>
testing <- train1[-split,] #test set of train data, which is validation set
#test1 is the test dataset
```

```
Data Exploration
library(dplyr)
library(purrr)
##
## Attaching package: 'purrr'
## The following object is masked from 'package:caret':
##
##
       lift
library(tidyr)
library(ggplot2)
#all variables - factor
training %>% keep(is.factor) %>% gather() %>%
  ggplot(aes(value)) +
  facet_wrap(~key, scales="free")+
  geom_bar()
## Warning: attributes are not identical across measure variables;
## they will be dropped
                Cabin
                                             Embarked
                                                                            Name
  500 -
                                400 -
                                                              300 -
  400 -
                                300 -
  300 -
                                                              200 -
                                200 -
```

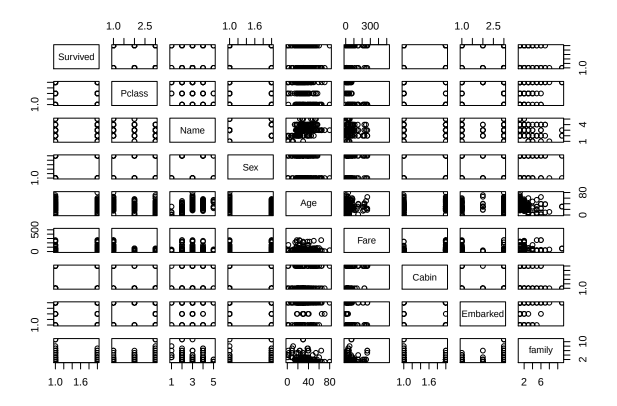


```
#all variables - factor
training %>% keep(is.numeric) %>% gather() %>%
ggplot(aes(value)) +
```

```
facet_wrap(~key, scales="free")+
geom_histogram(bins=30)+
geom_density()
```



pairs(training,col=train\$Survived)



## Logistic Regression

```
library(boot)
##
## Attaching package: 'boot'
## The following object is masked from 'package:lattice':
##
##
       melanoma
library(ROCR)
## Loading required package: gplots
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
##
       lowess
logistic.fit <- glm(Survived ~., data=training, family=binomial)</pre>
summary(logistic.fit)
##
## Call:
## glm(formula = Survived ~ ., family = binomial, data = training)
## Deviance Residuals:
##
       Min
                 1Q
                     Median
                                    3Q
                                            Max
```

```
## -2.3109 -0.5738 -0.4029 0.5231
                                       2.4922
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                4.052945
                           0.891576
                                     4.546 5.47e-06 ***
                           0.498261 -0.570 0.568856
## Pclass2
              -0.283878
## Pclass3
                           0.499055 -2.596 0.009440 **
               -1.295398
              -16.581174 639.499709 -0.026 0.979314
## NameMiss
## NameMr
               -3.749260
                           0.651305 -5.757 8.59e-09 ***
## NameMrs
              -15.786362 639.499776 -0.025 0.980306
## Nameunknown -3.735382
                           0.973263 -3.838 0.000124 ***
## Sex1
               15.702310 639.499405
                                      0.025 0.980411
## Age
               -0.019111
                           0.010537 -1.814 0.069724 .
                           0.003241
## Fare
                0.003516
                                     1.085 0.278094
## Cabin1
                0.860558
                           0.425625 2.022 0.043190 *
## EmbarkedQ
               -0.072022
                           0.462585 -0.156 0.876273
                           0.299850 -1.309 0.190403
## EmbarkedS
               -0.392620
## family
               -0.502213
                           0.107321 -4.680 2.88e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 832.49 on 624 degrees of freedom
## Residual deviance: 512.66 on 611 degrees of freedom
## AIC: 540.66
##
## Number of Fisher Scoring iterations: 14
logistic.probs <- predict(logistic.fit, newdata = testing, type="response")</pre>
logistic.pred <- ifelse(logistic.probs > 0.5, 1, 0)
test.pred <- testing$Survived</pre>
table(logistic.pred, test.pred)
##
                test.pred
## logistic.pred
                  0
                     1
               0 148 26
               1 16 76
sum(diag(table(logistic.pred, test.pred)))/nrow(testing)
## [1] 0.8421053
\#accuracy = 84.21\%
#cv error with 10 folds
cv.glm(training, logistic.fit, K=10)$delta[1]
## [1] 0.1336628
#around 13% error rate
ROCpred <- prediction(logistic.pred, test.pred)</pre>
```

```
ROCperf <- performance(ROCpred, "tpr", "fpr")</pre>
plot(ROCperf, colorize = TRUE, print.cutoffs.at = seq(0.1,0.1), text.adj = c(-0.2,1.7))
                                                                                0.1
      0.8
                                                                                             1.6
True positive rate
      9.0
      0.4
      0.2
                                                                                             4
                                                                                             o.
      0.0
                           0.2
             0.0
                                          0.4
                                                        0.6
                                                                      8.0
                                                                                     1.0
                                         False positive rate
performance(ROCpred, "auc")@y.values
## [[1]]
## [1] 0.8237685
#AUC value = 0.8237
confusionMatrix(as.factor(logistic.pred), test.pred)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
##
             0 148
                    26
             1 16
                   76
##
##
##
                   Accuracy : 0.8421
##
                     95% CI : (0.7926, 0.8838)
##
       No Information Rate: 0.6165
       P-Value [Acc > NIR] : 6.804e-16
##
##
##
                      Kappa: 0.6598
##
    Mcnemar's Test P-Value : 0.1649
##
##
                Sensitivity: 0.9024
##
               Specificity: 0.7451
##
##
             Pos Pred Value: 0.8506
             Neg Pred Value: 0.8261
##
                 Prevalence: 0.6165
##
```

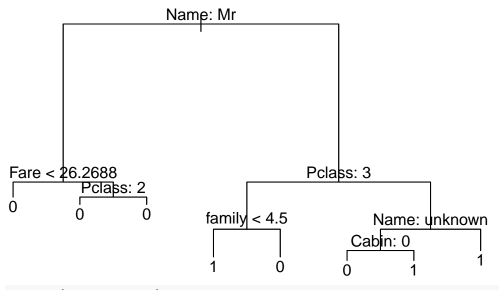
```
## Detection Rate : 0.5564
## Detection Prevalence : 0.6541
## Balanced Accuracy : 0.8238
##
## 'Positive' Class : 0
##
```

Logistic Regression Accuracy -> 84.21%

#### **Decision Tree**

```
library(tree)

tree.training <- tree(Survived~., training)
plot(tree.training);text(tree.training,pretty=0)</pre>
```

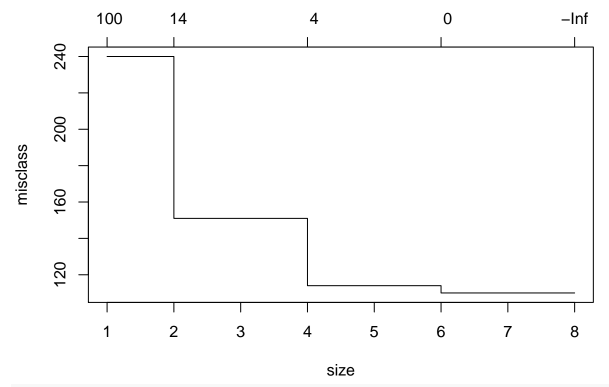


```
summary(tree.training)
```

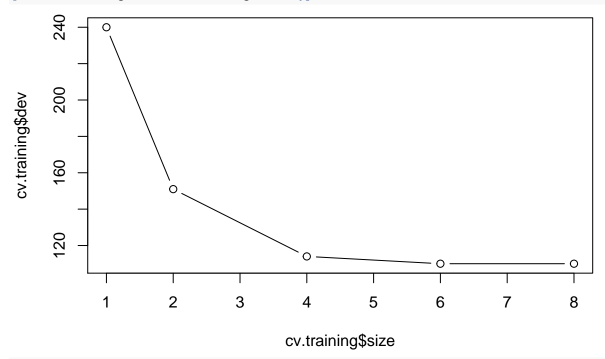
```
##
## Classification tree:
## tree(formula = Survived ~ ., data = training)
## Variables actually used in tree construction:
## [1] "Name" "Fare" "Pclass" "family" "Cabin"
## Number of terminal nodes: 8
## Residual mean deviance: 0.7791 = 480.7 / 617
## Misclassification error rate: 0.1632 = 102 / 625
tree.pred=predict(tree.training,testing,type="class") #with test datasets
table(tree.pred , testing$Survived)
```

```
## ## tree.pred 0 1 ## 0 147 28 ## 1 17 74
```

```
sum(diag(table(tree.pred, testing$Survived)))/nrow(testing)
## [1] 0.8308271
#accuracy 83.08%
deviance(tree.training)
## [1] 480.6808
misclass.tree(tree.training)
## [1] 102
#pruning via cv
cv.training=cv.tree(tree.training,FUN=prune.misclass)
cv.training
## $size
## [1] 8 6 4 2 1
##
## $dev
## [1] 110 110 114 151 240
## $k
## [1] -Inf 0.0 4.0 13.5 103.0
##
## $method
## [1] "misclass"
## attr(,"class")
## [1] "prune"
                       "tree.sequence"
plot(cv.training)
```

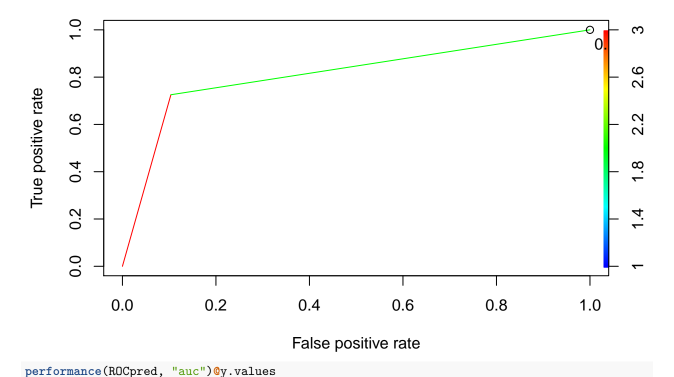


plot(cv.training\$size, cv.training\$dev, type="b")



prune.training <- prune.misclass(tree.training,best=7)
plot(prune.training);text(prune.training,pretty=0)</pre>

```
Name: Mr
                                            Pclass: 3
Fare < 26.2688
          Pclass: 2
0
                             family < 4.5
                                                     Name: unknown
                                                  Cabin: 0
                                       0
#pruned prediction
tree.pred <- predict(prune.training, testing, type="class")</pre>
table(tree.pred, testing$Survived)
## tree.pred 0
##
           0 147 28
           1 17 74
##
sum(diag(table(tree.pred, testing$Survived)))/nrow(testing)
## [1] 0.8308271
#83.08%
class(tree.pred)
## [1] "factor"
class(test.pred)
## [1] "factor"
ROCpred <- prediction(as.numeric(tree.pred), test.pred)</pre>
ROCperf <- performance(ROCpred, "tpr", "fpr")</pre>
plot(ROCperf, colorize = TRUE, print.cutoffs.at = seq(0.1,0.1), text.adj = c(-0.2,1.7))
```



```
## [[1]]
## [1] 0.8109158
#AUC = 0.8109
confusionMatrix(tree.pred, test.pred)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
                    1
##
            0 147
            1 17
                  74
##
##
                  Accuracy: 0.8308
##
##
                    95% CI: (0.7803, 0.8738)
       No Information Rate: 0.6165
##
##
       P-Value [Acc > NIR] : 2.211e-14
##
##
                     Kappa : 0.6348
##
##
    Mcnemar's Test P-Value : 0.136
##
               Sensitivity: 0.8963
##
##
               Specificity: 0.7255
            Pos Pred Value : 0.8400
##
##
            Neg Pred Value: 0.8132
                Prevalence: 0.6165
##
##
            Detection Rate: 0.5526
##
      Detection Prevalence: 0.6579
##
         Balanced Accuracy: 0.8109
```

#### Random Forest

```
library(randomForest)
## randomForest 4.6-14
## Type rfNews() to see new features/changes/bug fixes.
##
## Attaching package: 'randomForest'
## The following object is masked from 'package:dplyr':
##
##
       combine
## The following object is masked from 'package:ggplot2':
##
##
       margin
library (MASS)
#base random forest
rf.training <- randomForest(Survived~., data=training, importance=TRUE)
rf.training
##
## Call:
## randomForest(formula = Survived ~ ., data = training, importance = TRUE)
##
                  Type of random forest: classification
##
                        Number of trees: 500
## No. of variables tried at each split: 2
##
##
           OOB estimate of error rate: 17.44%
## Confusion matrix:
       0
          1 class.error
## 0 348 37 0.0961039
              0.3000000
## 1 72 168
#2 mtry - classification defaults = sqrt(p) = sqrt(8) ~= 2
#n.tree = 500, oob error rate = aroun 17%
#find best mtry
oob.err=double(8)
test.err=double(8)
for(mtry in 1:8){
  fit <- randomForest(Survived~., data=training, mtry=mtry, ntree=500)</pre>
  oob.err[mtry] <- 1-sum(diag(table(training$Survived,fit$predicted)))/nrow(training)</pre>
```

```
pred <- predict(fit, testing)</pre>
  test.err[mtry] <- 1-sum(diag(table(testing$Survived,pred)))/nrow(testing)</pre>
  cat(mtry," ")
     2 3 4 5 6 7 8
## 1
matplot(1:mtry,cbind(test.err,oob.err),pch=19,col=c("red","blue"),type="b",ylab="Error Rate")
legend("topright",legend=c("Test","00B"),pch=19,col=c("red","blue"))
      0.185
                                                                                Test
                                                                                OOB
Error Rate
      0.165
      0.155
                        2
              1
                                  3
                                            4
                                                      5
                                                                6
                                                                          7
                                                                                    8
                                              1:mtry
#2 mtry might be the best
#updated random forest
rf.training.update <- randomForest(Survived~., data=training, mtry=2, importance=TRUE, ntree=500)
rf.training.update
##
## Call:
    randomForest(formula = Survived ~ ., data = training, mtry = 2,
                                                                            importance = TRUE, ntree = 500
##
                  Type of random forest: classification
##
                         Number of trees: 500
##
## No. of variables tried at each split: 2
##
##
           OOB estimate of error rate: 17.44%
## Confusion matrix:
##
       0
           1 class.error
## 0 349 36 0.09350649
## 1 73 167 0.30416667
rf.pred <- predict(rf.training.update, testing)</pre>
table(testing$Survived, rf.pred)
```

rf.pred

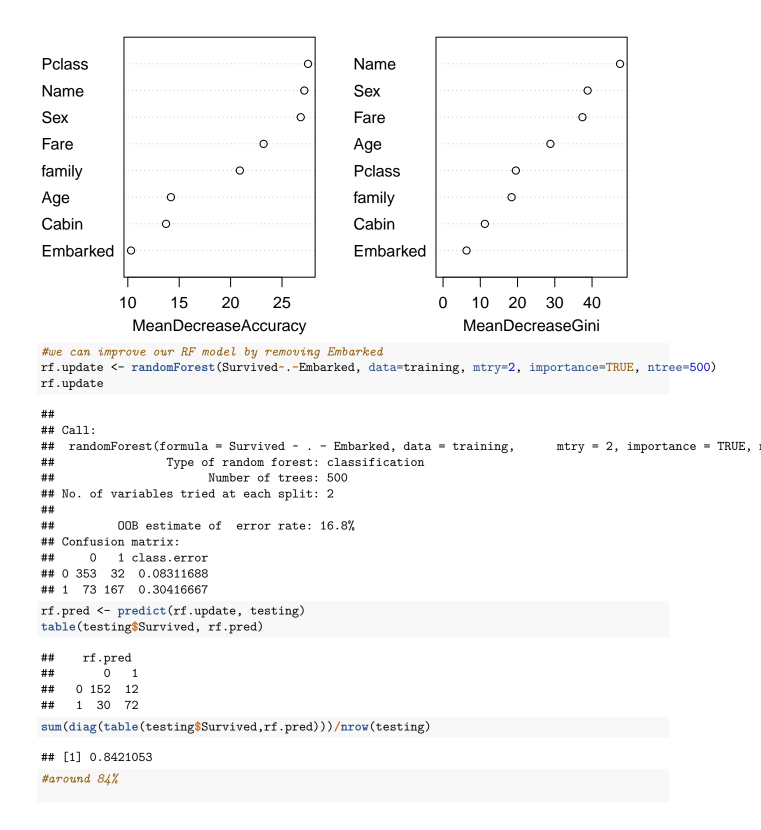
0

##

##

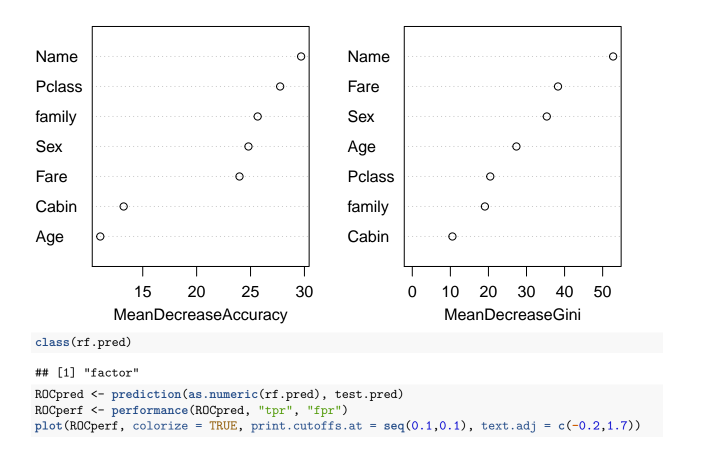
```
##
     0 153 11
##
    1 30 72
sum(diag(table(testing$Survived,rf.pred)))/nrow(testing)
## [1] 0.8458647
#around 84.58% accuracy
#since random forest provides different accuracy in every try,
#I made a function to get a mean value of the accuracy with few number of tries
mean.acc <- function(training, testing, mtry, ntree, try, formula){</pre>
  acc <- NULL
 for(i in 1:try){
  randomFtrain <- randomForest(formula, data=training, mtry=mtry, importance=TRUE, ntree=ntree)</pre>
  randomFpred <- predict(randomFtrain, testing)</pre>
  acc[i] <- sum(diag(table(testing$Survived, randomFpred)))/nrow(testing)</pre>
  }
  return(mean(acc))
#2 mtry, 500 ntree, 10 tries
mean.acc(training = training, testing = testing, mtry = 2, ntree = 500, try = 10, formula = as.formula(
## [1] 0.8454887
#around 84.5% accuracy
importance(rf.training.update)
                              1 MeanDecreaseAccuracy MeanDecreaseGini
##
## Pclass 18.134499 20.011172
                                            27.53240
                                                            19.545816
## Name
           23.064960 20.999153
                                            27.16987
                                                            47.525815
## Sex
           23.792787 20.190242
                                                            38.822613
                                           26.82020
## Age
           11.570976 7.040736
                                           14.19494
                                                            28.820272
           14.556035 15.535657
                                            23.20695
                                                            37.421976
## Fare
## Cabin
           12.280560 1.776714
                                            13.72102
                                                            11.226495
                                            10.31469
## Embarked 6.638619 7.246604
                                                            6.311426
## family
           21.015530 5.218679
                                            20.89440
                                                            18.407316
varImpPlot(rf.training.update)
```

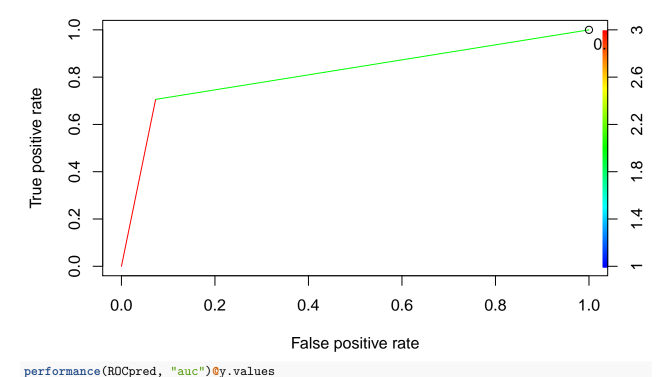
## rf.training.update



```
mean.acc(training, testing, mtry = 2, ntree = 500, try = 10, formula = as.formula(Survived~.-Embarked))
## [1] 0.8421053
#accruacy around 84.3%
importance(rf.update)
                            1 MeanDecreaseAccuracy MeanDecreaseGini
## Pclass 19.281870 20.462255
                                          27.74701
                                                            20.47984
          26.192644 23.155114
                                          29.69098
                                                            52.75031
          22.439350 19.452844
                                          24.80557
                                                            35.32776
## Sex
## Age
           7.367302 8.111400
                                          11.06260
                                                            27.34985
## Fare
                                          23.97459
          14.755373 17.864728
                                                            38.27450
## Cabin 12.514360 1.968030
                                          13.22990
                                                            10.53989
## family 24.767579 7.058626
                                          25.66010
                                                            19.07239
varImpPlot(rf.update)
```

# rf.update





```
## [[1]]
## [1] 0.8163558
#AUC ~= around 0.8212
confusionMatrix(rf.pred, test.pred)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
                    1
##
            0 152
                   30
            1 12 72
##
##
##
                  Accuracy : 0.8421
##
                    95% CI: (0.7926, 0.8838)
       No Information Rate: 0.6165
##
##
       P-Value [Acc > NIR] : 6.804e-16
##
##
                     Kappa : 0.6545
##
##
    Mcnemar's Test P-Value : 0.008712
##
               Sensitivity: 0.9268
##
##
               Specificity: 0.7059
            Pos Pred Value : 0.8352
##
##
            Neg Pred Value: 0.8571
                Prevalence: 0.6165
##
##
            Detection Rate: 0.5714
##
      Detection Prevalence: 0.6842
##
         Balanced Accuracy: 0.8164
```

```
##
##
           'Positive' Class: 0
##
Random Forest Accuracy -> around 84.21\%
Boosting
library(gbm)
## Loaded gbm 2.1.5
set.seed(123)
boost.training <- gbm(as.character(Survived)~., data=training,</pre>
                       distribution = "bernoulli", n.trees = 500,
                       interaction.depth = 3, shrinkage = 0.01, cv.folds=5)
summary(boost.training)
Name
Pclass
Age
Embarked
     0
                      10
                                       20
                                                        30
                                                                         40
                                  Relative influence
##
                        rel.inf
                  var
## Name
                 Name 45.234382
## Fare
                 Fare 14.554858
               Pclass 13.760218
## Pclass
```

```
table(training$Survived, boost.pred)
##
      boost.pred
##
         0
           1
##
     0 356 29
##
     1 54 186
sum(diag(table(training$Survived, boost.pred)))/nrow(training)
## [1] 0.8672
#86.08% accuracy
#test on the test set
boost.pred <- predict(boost.training, newdata = testing, n.trees = 500, type = "response")
boost.pred <- ifelse(boost.pred > 0.5, 1, 0)
table(testing$Survived, boost.pred)
##
      boost.pred
##
         0
##
     0 148 16
     1 29 73
sum(diag(table(testing$Survived, boost.pred)))/nrow(testing)
## [1] 0.8308271
#83.83% accuracy
#cross validation to get the best ntree
folds <- sample(rep(1:5, length = nrow(training)))</pre>
folds
     [1] \ 4 \ 2 \ 1 \ 3 \ 3 \ 2 \ 4 \ 2 \ 4 \ 3 \ 3 \ 1 \ 2 \ 1 \ 2 \ 3 \ 2 \ 5 \ 1 \ 3 \ 2 \ 5 \ 1 \ 3 \ 1 \ 5 \ 4 \ 2 \ 2 \ 3 \ 5 \ 1 \ 4
##
  [36] 5 2 2 5 2 1 4 1 3 4 3 1 3 5 4 2 5 2 3 1 1 1 1 3 5 1 4 5 4 1 4 1 4 1 1
## [71] 5 5 3 2 2 2 3 1 2 5 2 1 2 1 4 5 2 5 4 5 1 4 2 5 3 4 3 1 5 1 2 3 1 4 1
## [106] 3 2 5 3 3 2 5 4 5 5 5 5 5 5 3 3 2 3 2 4 5 1 4 1 5 1 5 4 3 5 1 3 3 5
## [141] 4 5 2 5 3 3 1 4 1 1 2 2 4 5 1 2 3 1 3 2 4 5 1 3 4 5 5 5 5 1 3 3 1 4 5 1
## [176] 4 1 4 1 5 4 2 1 5 1 2 3 3 2 5 4 2 2 2 4 1 3 5 3 5 2 1 5 1 1 4 4 5 2 4
## [211] 4 4 4 2 1 4 2 5 4 4 4 3 1 5 3 1 4 1 1 5 4 2 5 3 3 2 2 5 4 2 3 1 2 5 1
## [246] 2 3 4 3 4 4 5 5 5 2 3 2 4 3 1 5 3 1 4 5 5 2 1 3 2 3 3 3 4 2 5 5 2 4 5
## [281] 5 3 2 2 1 4 5 2 1 4 1 3 5 1 5 1 4 3 3 3 4 3 5 4 4 1 1 1 3 3 2 5 4 4 1
## [316] 5 3 3 4 4 2 4 1 4 3 1 3 3 3 5 4 5 4 3 1 3 4 4 1 4 1 5 4 2 2 5 1 5 4 3
## [351] 3 3 5 2 2 5 3 1 3 5 4 1 1 3 5 3 3 2 2 2 3 5 4 4 2 3 2 5 4 2 3 1 1 2 4
## [386] 2 5 5 3 4 4 4 1 2 5 2 3 5 1 2 1 3 3 1 4 2 5 1 2 3 4 4 3 1 3 2 3 2 3 5
## [421] 1 2 3 4 1 1 1 5 1 3 1 2 5 3 5 3 1 2 2 3 5 2 4 5 2 3 1 3 4 2 3 2 3 1 1
## [456] 5 1 4 4 4 3 4 1 2 2 5 2 1 4 3 1 3 1 5 4 5 5 4 2 2 2 2 4 5 5 5 2 3 5 3
## [491] 3 4 3 1 2 2 4 4 5 4 5 2 1 2 3 4 2 3 1 2 1 2 5 1 3 2 2 1 4 5 3 2 2 4 5
## [526] 2 4 5 2 5 2 1 5 5 1 1 2 4 5 2 4 4 5 1 3 1 4 2 5 1 4 4 3 3 3 4 1 1 1 1 4
## [561] 5 2 2 1 4 1 1 5 4 3 4 3 4 5 2 3 3 2 4 3 1 1 2 5 2 3 5 5 4 5 4 1 4 5 1
## [596] 5 4 4 4 1 2 1 1 5 2 3 4 3 4 5 2 2 4 2 5 2 5 3 3 5 4 5 2 3 4
```

```
table(folds)
## folds
## 1 2
             3 4
                     5
## 125 125 125 125 125
ntree \leftarrow seq(500, 4000, by=500)
cv.errors <- matrix(NA, length(ntree),5)</pre>
cv.errors <- cbind(ntree, cv.errors)</pre>
cv.errors
##
        ntree
## [1,]
        500 NA NA NA NA NA
## [2,] 1000 NA NA NA NA NA
## [3,] 1500 NA NA NA NA NA
## [4,] 2000 NA NA NA NA NA
## [5,] 2500 NA NA NA NA NA
## [6,] 3000 NA NA NA NA NA
## [7,] 3500 NA NA NA NA NA
## [8,] 4000 NA NA NA NA NA
for(i in 1:5){
  for(j in ntree){
    boost.fit <- gbm(as.character(Survived)~.,</pre>
                     data = training[folds != i,], distribution = "bernoulli",
                     n.trees = j, interaction.depth = 3,
                     shrinkage = 0.01, verbose = F)
    boost.pred <- predict(boost.fit, training[folds == i,], n.trees = j)</pre>
    boost.pred <- ifelse(boost.pred > 0.5, 1, 0)
    cv.errors[which(j==ntree), i+1] =
      1- sum(diag(table(training$Survived[folds==i], boost.pred)))/nrow(training[folds==i,])
  }
}
cv.errors
##
        ntree
## [1,]
        500 0.240 0.144 0.200 0.184 0.152
## [2,] 1000 0.232 0.128 0.208 0.176 0.160
## [3,] 1500 0.248 0.136 0.208 0.160 0.168
## [4,] 2000 0.232 0.136 0.208 0.160 0.168
## [5,] 2500 0.256 0.144 0.216 0.160 0.160
## [6,] 3000 0.248 0.128 0.216 0.168 0.144
## [7,] 3500 0.256 0.136 0.232 0.176 0.152
## [8,] 4000 0.256 0.144 0.224 0.176 0.168
cv.errors1 <- cv.errors[,-1]
cv.errors1
##
## [1,] 0.240 0.144 0.200 0.184 0.152
## [2,] 0.232 0.128 0.208 0.176 0.160
## [3,] 0.248 0.136 0.208 0.160 0.168
## [4,] 0.232 0.136 0.208 0.160 0.168
```

```
## [5,] 0.256 0.144 0.216 0.160 0.160

## [6,] 0.248 0.128 0.216 0.168 0.144

## [7,] 0.256 0.136 0.232 0.176 0.152

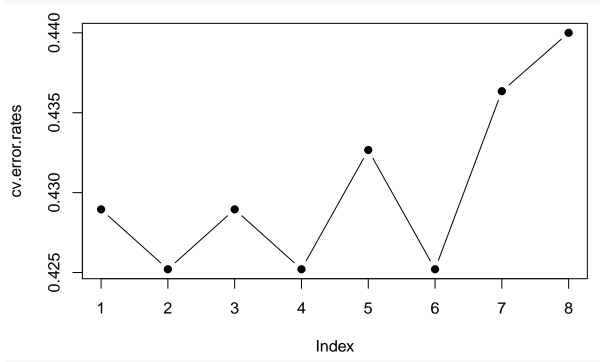
## [8,] 0.256 0.144 0.224 0.176 0.168

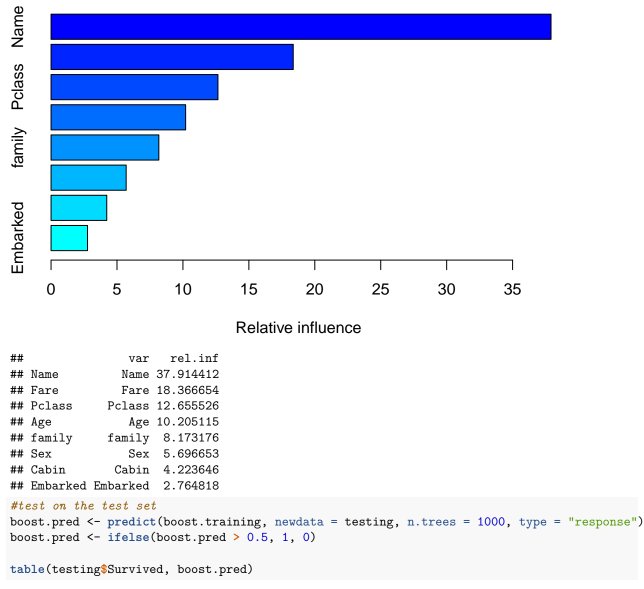
cv.error.rates=sqrt(apply(cv.errors1,1,mean))

which.min(cv.error.rates)
```

#### ## [1] 2

plot(cv.error.rates,pch=19,type="b")





```
table(testing$Survived, boost.pred)

## boost.pred
## 0 1
## 0 147 17
```

sum(diag(table(testing\$Survived, boost.pred)))/nrow(testing)

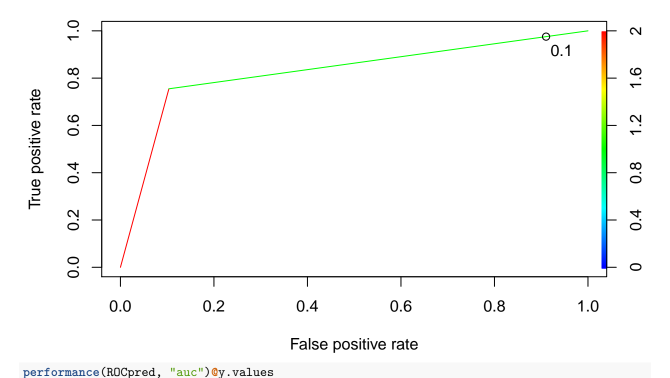
```
## [1] 0.8421053
```

##

1 25 77

```
#84.21% accuracy

ROCpred <- prediction(as.numeric(boost.pred), test.pred)
ROCperf <- performance(ROCpred, "tpr", "fpr")
plot(ROCperf, colorize = TRUE, print.cutoffs.at = seq(0.1,0.1), text.adj = c(-0.2,1.7))</pre>
```



```
## [[1]]
## [1] 0.8256217

#auc = 0.8256

confusionMatrix(as.factor(boost.pred), testing$Survived)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
                0
                    1
##
            0 147
                   25
            1 17
                  77
##
##
                  Accuracy : 0.8421
##
##
                    95% CI: (0.7926, 0.8838)
       No Information Rate: 0.6165
##
##
       P-Value [Acc > NIR] : 6.804e-16
##
##
                     Kappa: 0.661
##
##
    Mcnemar's Test P-Value : 0.2801
##
               Sensitivity: 0.8963
##
##
               Specificity: 0.7549
            Pos Pred Value : 0.8547
##
##
            Neg Pred Value: 0.8191
##
                Prevalence: 0.6165
##
            Detection Rate: 0.5526
##
      Detection Prevalence: 0.6466
##
         Balanced Accuracy: 0.8256
```

```
##
##
          'Positive' Class: 0
##
Boosting -> 84.21\%
SVM - linear
library(e1071)
set.seed(123)
svm.fit <- svm(Survived~., data=training, scale=FALSE, kernel="linear", cost=5)</pre>
print(svm.fit)
##
## Call:
## svm(formula = Survived ~ ., data = training, kernel = "linear",
       cost = 5, scale = FALSE)
##
##
##
## Parameters:
##
      SVM-Type: C-classification
## SVM-Kernel: linear
##
          cost: 5
##
         gamma: 0.07142857
##
## Number of Support Vectors: 258
summary(svm.fit)
##
## Call:
## svm(formula = Survived ~ ., data = training, kernel = "linear",
##
       cost = 5, scale = FALSE)
##
```

```
##
## Parameters:
##
      SVM-Type: C-classification
   SVM-Kernel: linear
         cost: 5
##
##
         gamma: 0.07142857
##
## Number of Support Vectors: 258
##
## ( 130 128 )
##
## Number of Classes: 2
## Levels:
## 0 1
tune.out <- tune(svm, Survived~., data=training,</pre>
```

kernel = "linear",

ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10,15,20)))

```
summary(tune.out)
## Parameter tuning of 'svm':
## - sampling method: 10-fold cross validation
##
## - best parameters:
## cost
##
      10
##
## - best performance: 0.1646953
##
## - Detailed performance results:
##
               error dispersion
       cost
## 1 0.001 0.3808244 0.04926884
## 2 0.010 0.2030978 0.04264397
## 3 0.100 0.1967486 0.04079913
## 4 1.000 0.1710701 0.03838288
## 5 5.000 0.1695341 0.02972506
## 6 10.000 0.1646953 0.03213910
## 7 15.000 0.1646953 0.03213910
## 8 20.000 0.1646953 0.03213910
#best model with the C values (tuning parameter)
tune.out$best.model
##
## Call:
## best.tune(method = svm, train.x = Survived ~ ., data = training,
       ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10, 15, 20)),
##
       kernel = "linear")
##
##
## Parameters:
      SVM-Type: C-classification
##
##
  SVM-Kernel: linear
##
          cost: 10
##
         gamma: 0.07142857
##
## Number of Support Vectors: 255
svm.best <- tune.out$best.model</pre>
summary(svm.best)
##
## Call:
## best.tune(method = svm, train.x = Survived ~ ., data = training,
       ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10, 15, 20)),
       kernel = "linear")
##
##
##
## Parameters:
##
      SVM-Type: C-classification
## SVM-Kernel: linear
```

```
##
           cost:
                  10
         gamma: 0.07142857
##
##
## Number of Support Vectors:
##
##
    (129 126)
##
##
## Number of Classes: 2
##
## Levels:
   0 1
##
svm.pred <- predict(svm.best, testing, type="class")</pre>
table(predict = svm.pred, truth = testing$Survived)
##
           truth
## predict
              0
                  1
         0 148
                28
##
         1
            16
                74
sum(diag(table(svm.pred, testing$Survived)))/nrow(testing)
## [1] 0.8345865
#83.46 accuracy
ROCpred <- prediction(as.numeric(svm.pred), test.pred)</pre>
ROCperf <- performance(ROCpred, "tpr", "fpr")</pre>
plot(ROCperf, colorize = TRUE, print.cutoffs.at = seq(0.1,0.1), text.adj = c(-0.2,1.7))
                                                                                       0
                                                                                        0
      0.8
                                                                                              9
True positive rate
      9.0
      0.4
      0.2
      0.0
             0.0
                            0.2
                                          0.4
                                                         0.6
                                                                       8.0
                                                                                      1.0
                                         False positive rate
performance(ROCpred, "auc")@y.values
```

## [[1]]

```
## [1] 0.8139646
\#auc = 0.8139
confusionMatrix(svm.pred, testing$Survived)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
               0
                  1
##
            0 148 28
            1 16 74
##
##
##
                  Accuracy : 0.8346
##
                    95% CI : (0.7844, 0.8772)
##
       No Information Rate: 0.6165
       P-Value [Acc > NIR] : 7.118e-15
##
##
##
                     Kappa: 0.6422
##
##
   Mcnemar's Test P-Value: 0.09725
##
               Sensitivity: 0.9024
##
##
               Specificity: 0.7255
##
            Pos Pred Value: 0.8409
##
            Neg Pred Value: 0.8222
                Prevalence: 0.6165
##
##
            Detection Rate: 0.5564
##
      Detection Prevalence: 0.6617
         Balanced Accuracy: 0.8140
##
##
##
          'Positive' Class : 0
##
SVM linear Accuracy -> 83.46%
SVM - radial kernel
svm.fit <- svm(Survived~., data=training, scale=FALSE, kernel="radial", cost=5)</pre>
print(svm.fit)
##
## Call:
## svm(formula = Survived ~ ., data = training, kernel = "radial",
       cost = 5, scale = FALSE)
##
##
##
## Parameters:
      SVM-Type: C-classification
##
    SVM-Kernel: radial
##
          cost:
```

##

##

gamma: 0.07142857

## Number of Support Vectors: 445

```
summary(svm.fit)
##
## Call:
## svm(formula = Survived ~ ., data = training, kernel = "radial",
       cost = 5, scale = FALSE)
##
##
## Parameters:
##
     SVM-Type: C-classification
## SVM-Kernel: radial
##
        cost: 5
##
        gamma: 0.07142857
##
## Number of Support Vectors: 445
##
  (215 230)
##
##
## Number of Classes: 2
##
## Levels:
## 0 1
tune.out <- tune(svm, Survived~., data=training,
                 kernel = "radial",
                 ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10, 15, 20)))
summary(tune.out)
## Parameter tuning of 'svm':
##
## - sampling method: 10-fold cross validation
## - best parameters:
## cost
##
       1
##
## - best performance: 0.1664363
## - Detailed performance results:
##
      cost
               error dispersion
## 1 0.001 0.3841014 0.06229791
## 2 0.010 0.3841014 0.06229791
## 3 0.100 0.1935996 0.05389965
## 4 1.000 0.1664363 0.04209901
## 5 5.000 0.1711982 0.04505016
## 6 10.000 0.1711214 0.04533815
## 7 15.000 0.1790835 0.05147086
## 8 20.000 0.1742960 0.05290177
tune.out$best.model
```

##

```
## Call:
## best.tune(method = svm, train.x = Survived ~ ., data = training,
       ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10, 15, 20)),
##
       kernel = "radial")
##
##
## Parameters:
      SVM-Type: C-classification
##
##
   SVM-Kernel: radial
##
          cost: 1
##
         gamma: 0.07142857
##
## Number of Support Vectors:
svm.best <- tune.out$best.model</pre>
summary(svm.best)
##
## Call:
## best.tune(method = svm, train.x = Survived ~ ., data = training,
       ranges = list(cost = c(0.001, 0.01, 0.1, 1, 5, 10, 15, 20)),
       kernel = "radial")
##
##
##
## Parameters:
      SVM-Type: C-classification
##
##
  SVM-Kernel: radial
##
         cost: 1
##
         gamma: 0.07142857
##
## Number of Support Vectors: 299
##
   ( 146 153 )
##
##
##
## Number of Classes: 2
##
## Levels:
svm.pred <- predict(svm.best, testing, type="class")</pre>
table(predict = svm.pred, truth = testing$Survived)
##
          truth
## predict
           0 1
         0 149 28
         1 15 74
sum(diag(table(svm.pred, testing$Survived)))/nrow(testing)
## [1] 0.8383459
#83.83% accuracy
ROCpred <- prediction(as.numeric(svm.pred), test.pred)</pre>
ROCperf <- performance(ROCpred, "tpr", "fpr")</pre>
```

```
plot(ROCperf, colorize = TRUE, print.cutoffs.at = seq(0.1,0.1), text.adj = c(-0.2,1.7))
                                                                                      0
                                                                                             က
                                                                                        0
      \infty
                                                                                              9
      o.
                                                                                              ď
True positive rate
      9.0
                                                                                              \sim
      0.4
                                                                                              \infty
      0.2
      0.0
             0.0
                           0.2
                                          0.4
                                                         0.6
                                                                       8.0
                                                                                      1.0
                                         False positive rate
performance(ROCpred, "auc")@y.values
## [[1]]
## [1] 0.8170134
\#auc = 0.8170
confusionMatrix(svm.pred, testing$Survived)
## Confusion Matrix and Statistics
##
##
              Reference
## Prediction
                 0
                     1
##
             0 149
                    28
##
             1 15
                   74
##
                   Accuracy : 0.8383
##
##
                     95% CI : (0.7885, 0.8805)
##
       No Information Rate: 0.6165
##
       P-Value [Acc > NIR] : 2.231e-15
##
##
                      Kappa: 0.6497
##
##
    Mcnemar's Test P-Value : 0.06725
##
                Sensitivity: 0.9085
##
                Specificity: 0.7255
##
            Pos Pred Value : 0.8418
##
##
             Neg Pred Value: 0.8315
                 Prevalence: 0.6165
##
##
             Detection Rate: 0.5602
```

```
## Detection Prevalence: 0.6654
## Balanced Accuracy: 0.8170
##

## 'Positive' Class: 0
##

SVM - radial Accuracy -> 83.83%

Logistic Regression Accuracy -> 84.21%

Decision Tree Accuracy -> 83.08%

Random Forest Accuracy -> around 84.5%

Boosting Accuracy -> 84.21%

SVM linear Accuracy -> 83.46%

SVM radial Accuracy -> 83.83%
```

## Creating submission file with RF model

```
final.pred <- predict(rf.update, newdata = test1, type="response")

final <- data.frame(PassengerId = test$PassengerId, FinalPred = final.pred)
head(final)</pre>
```

```
##
       PassengerId FinalPred
## 892
               892
## 893
               893
                           1
## 894
               894
                           0
               895
                           0
## 895
## 896
               896
                           1
## 897
               897
                           0
```