## **Assignment 2**

# Predicting the type of a Breast Tumor (benign or malignant) Using Random-Forest Model

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4/12/2020

**The data is loaded using the mlbench library, data(BreastCancer)** A data frame with 699 observations on 11 variables, one being a character variable, 9 being ordered or nominal, and 1 target class.

```
library(caret)
## Loading required package: lattice
## Loading required package: ggplot2
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(ggplot2)
```

\*\*\* Understanding Data structure

```
summary(BreastCancer)
##
         Ιd
                         Cl.thickness
                                         Cell.size
                                                        Cell.shape
                                                                     Marg.adhesio
n
##
    Length:699
                        1
                                :145
                                       1
                                               :384
                                                      1
                                                              :353
                                                                     1
                                                                             :407
    Class :character
                        5
                                                              : 59
                                                                     2
                                                                             : 58
##
                                :130
                                       10
                                               : 67
                                                      2
    Mode :character
                        3
                                                                             : 58
##
                                :108
                                       3
                                               : 52
                                                      10
                                                              : 58
                                                                     3
##
                        4
                                : 80
                                       2
                                               : 45
                                                      3
                                                              : 56
                                                                     10
                                                                             : 55
##
                        10
                                : 69
                                       4
                                               : 40
                                                      4
                                                              : 44
                                                                     4
                                                                             : 33
                                       5
                                               : 30
                                                      5
                                                              : 34
                                                                             : 25
##
                                : 50
                                                                     8
                                       (Other): 81
                                                                     (Other): 63
##
                        (Other):117
                                                      (Other): 95
##
     Epith.c.size Bare.nuclei
                                   Bl.cromatin
                                                Normal.nucleoli
                                                                     Mitoses
                         :402
          :386
                                  2
                                         :166
                                                 1
                                                        :443
                                                                 1
                                                                         :579
##
                   1
```

```
## 3
           : 72
                  10
                          :132
                                               10
                                                       : 61
                                 3
                                        :165
                                                                        : 35
   4
           : 48
                          : 30
                                        :152
                                                       : 44
                                                                3
                                                                       : 33
##
                  2
                                 1
                                               3
                                 7
                                        : 73
                                                       : 36
                                                                10
                                                                       : 14
##
   1
           : 47
                  5
                          : 30
                                               2
                                                                       : 12
## 6
           : 41
                          : 28
                                 4
                                        : 40
                                               8
                                                       : 24
                                                                4
                  3
                                 5
                                        : 34
                                                                7
## 5
           : 39
                  (Other): 61
                                               6
                                                       : 22
                                                                       : 9
    (Other): 66
                  NA's
                        : 16
                                 (Other): 69
                                               (Other): 69
                                                                (Other): 17
##
##
          Class
##
    benign
             :458
    malignant:241
##
##
##
##
##
str(BreastCancer)
## 'data.frame':
                    699 obs. of 11 variables:
                      : chr "1000025" "1002945" "1015425" "1016277"
## $ Id
## $ Cl.thickness
                      : Ord.factor w/ 10 levels "1"<"2"<"3"<"4"<...: 5 5 3 6 4
8 1 2 2 4 ...
                      : Ord.factor w/ 10 levels "1"<"2"<"3"<"4"<..: 1 4 1 8 1
## $ Cell.size
10 1 1 1 2 ...
                      : Ord.factor w/ 10 levels "1"<"2"<"3"<"4"<...: 1 4 1 8 1
## $ Cell.shape
10 1 2 1 1 ...
## $ Marg.adhesion : Ord.factor w/ 10 levels "1"<"2"<"3"<"4"<..: 1 5 1 1 3
8 1 1 1 1 ...
                      : Ord.factor w/ 10 levels "1"<"2"<"3"<"4"<..: 2 7 2 3 2
## $ Epith.c.size
7 2 2 2 2 ...
## $ Bare.nuclei
                      : Factor w/ 10 levels "1", "2", "3", "4", ...: 1 10 2 4 1 10
10 1 1 1 ...
## $ Bl.cromatin
                      : Factor w/ 10 levels "1", "2", "3", "4", ...: 3 3 3 3 3 9 3
3 1 2 ...
## $ Normal.nucleoli: Factor w/ 10 levels "1", "2", "3", "4",..: 1 2 1 7 1 7 1
1 1 1 ...
                      : Factor w/ 9 levels "1", "2", "3", "4", ...: 1 1 1 1 1 1 1 1 1
## $ Mitoses
5 1 ...
## $ Class
                     : Factor w/ 2 levels "benign", "malignant": 1 1 1 1 1 2 1
1 1 1 ...
levels(BreastCancer$Class)
## [1] "benign"
                   "malignant"
```

Let's calculate the number and percent of missing data and plot them \*\* checking if there any missing data using "Amelia package"

```
# Check if there are any missing values:
anyNA(BreastCancer)
## [1] TRUE
```

```
sum(is.na(BreastCancer))
## [1] 16
```

#### Ploting the missing and observed data values

```
library(Amelia)

## Loading required package: Rcpp

## ##

## ## Amelia II: Multiple Imputation

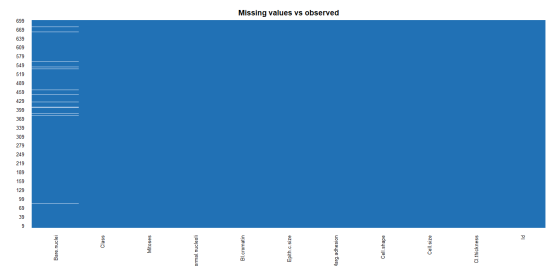
## ## (Version 1.7.6, built: 2019-11-24)

## ## Copyright (C) 2005-2020 James Honaker, Gary King and Matthew Blackwell

## ## Refer to http://gking.harvard.edu/amelia/ for more information

## ##

missmap(BreastCancer, main = "Missing values vs observed",legend = FALSE)
```



```
mean(is.na(BreastCancer))
## [1] 0.002080895
```

\*\* we have 16 missing values in our dataset.

#### **Cleaning missing data**

```
Breast <- na.omit(BreastCancer)[,c(2:11)]

library(randomForest)

## randomForest 4.6-14

## Type rfNews() to see new features/changes/bug fixes.

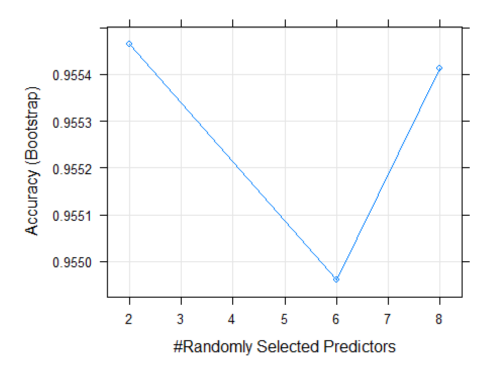
##

## Attaching package: 'randomForest'</pre>
```

```
## The following object is masked from 'package:dplyr':
##
##
       combine
## The following object is masked from 'package:ggplot2':
##
       margin
set.seed(123)
intrain <- createDataPartition(y = Breast$Class, p= 0.7, list = FALSE)
training <- Breast[intrain,]</pre>
testing <- Breast[-intrain,]</pre>
set.seed(123)
rf.model<-train(Class~.,data=training,method='rf')</pre>
rf.model
## Random Forest
##
## 479 samples
     9 predictor
     2 classes: 'benign', 'malignant'
##
##
## No pre-processing
## Resampling: Bootstrapped (25 reps)
## Summary of sample sizes: 479, 479, 479, 479, 479, ...
## Resampling results across tuning parameters:
##
##
     mtry Accuracy
                      Kappa
##
     2
           0.9554657 0.9033081
           0.9499567 0.8908242
##
     41
##
     80
           0.9455779 0.8813470
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
```

### **Grid search with Bootstrapped Resampling**

```
##
## No pre-processing
## Resampling: Bootstrapped (25 reps)
## Summary of sample sizes: 479, 479, 479, 479, 479, 479, ...
## Resampling results across tuning parameters:
##
##
     mtry
          Accuracy
                      Kappa
##
     2
           0.9554657
                      0.9033081
##
     6
           0.9549596
                      0.9021462
     8
           0.9554124 0.9031210
##
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
plot(RF_Grid_Boot)
```



```
preds_rf_boot <- predict(RF_Grid_Boot, testing[1:9])

confusionMatrix(table(preds_rf_boot, testing$Class))

## Confusion Matrix and Statistics
##

##

## preds_rf_boot benign malignant
## benign 129 3

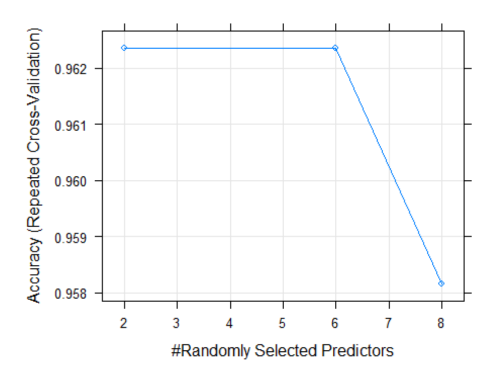
## malignant 4 68
##</pre>
```

```
##
                  Accuracy : 0.9657
##
                    95% CI: (0.9306, 0.9861)
       No Information Rate : 0.652
##
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.9246
##
##
   Mcnemar's Test P-Value : 1
##
##
               Sensitivity: 0.9699
##
               Specificity: 0.9577
##
            Pos Pred Value: 0.9773
##
            Neg Pred Value: 0.9444
##
                Prevalence: 0.6520
##
            Detection Rate: 0.6324
##
      Detection Prevalence: 0.6471
##
         Balanced Accuracy: 0.9638
##
##
          'Positive' Class : benign
##
```

#### **Grid Search with Cross-Validation (10 fold, repeated 4 times)**

```
set.seed(123)
control <- trainControl(method="repeatedcv", number=10, repeats=3, search="gr</pre>
id")
Grid_Serach <- expand.grid(.mtry=c(2,6,8))</pre>
# Random forest Model Building
RF Grid_CV<-train(Class~.,</pre>
                 data=training,
                 method='rf',
                 tuneGrid=Grid Serach,
                 trControl=control
print(RF Grid CV)
## Random Forest
##
## 479 samples
##
     9 predictor
##
     2 classes: 'benign', 'malignant'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 430, 431, 431, 431, 432, 431, ...
## Resampling results across tuning parameters:
##
##
     mtry
           Accuracy
                       Kappa
           0.9623498 0.9179821
##
     2
##
     6
           0.9623646 0.9185535
##
     8
           0.9581536 0.9091281
```

```
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 6.
plot(RF_Grid_CV)
```



```
#Prediction using test data
preds_rf_cv <- predict(RF_Grid_CV, testing[1:9])</pre>
confusionMatrix(table(preds_rf_cv, testing$Class))
## Confusion Matrix and Statistics
##
##
## preds_rf_cv benign malignant
     benign
##
                  130
                               3
##
     malignant
                    3
                              68
##
##
                  Accuracy : 0.9706
                    95% CI: (0.9371, 0.9891)
##
##
       No Information Rate: 0.652
       P-Value [Acc > NIR] : <2e-16
##
##
##
                      Kappa: 0.9352
##
##
    Mcnemar's Test P-Value : 1
##
##
               Sensitivity: 0.9774
```

```
##
               Specificity: 0.9577
##
            Pos Pred Value : 0.9774
            Neg Pred Value : 0.9577
##
##
                Prevalence: 0.6520
            Detection Rate : 0.6373
##
##
      Detection Prevalence : 0.6520
         Balanced Accuracy : 0.9676
##
##
##
          'Positive' Class : benign
##
```

From the above analysis we can notice that The 10-fold cross validation has a better accuracy than bootstrapped resampling.