Heart Disease predicting model

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Contents

Н	arvardX: PH125.9	1
	Data Science Professional Certificate: Capstone - "Choose Your Own Project Submission"	1
	Introduction	1
	Methods	2
	cleaning and process the Dataset	2
	Data Exploration and testing correlations	5
	Creating the Training and Testing Sets	11
	Model 1: K-Nearest Neighbors	12
	Model 2: Adaptive boosting Model	14
	Model 3: General Logistic Regression model	15
	Model 4: Decision Tree model	16
	Model 5: Random Forest model	17
	Ensemble model	18
	Conclusions	19

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You can find this project on Github Website at: https://github.com/davidemori/Cardiac-Disease

Introduction

Of the 56.9 million deaths worldwide in 2016, more than half (54%) were due to the top 10 causes. Ischaemic heart disease and stroke are the world's biggest killers, accounting for a combined 15.2 million deaths in 2016. These diseases have remained the leading causes of death globally in the last 15 years (World Health Organization, 2019). The aim of this project is to analyze the causes and try to find a model to predict, on the basis of some parameters, the occurrence of an heart disease.

Many datasets about Medical Sciences, and in particular about heart diseases, are free downloadable on the "WEB". For the following project we have used a subset of the "Cleveland Heart Disease Dataset" from the UCI Archives, that can be found at the following link: http://archive.ics.uci.edu/ml/datasets/heart+disease

On this dataset we'll try to perform some exploratory analyses, and test a total of six models (5 + 1) ensemble model to find the best fit for prediction of the heart disease in an adult population.

Methods

cleaning and process the Dataset

The first goal of our cleaning and tidying process is to download the dataset and define the names of the columns. In fact, despite the dataset is provided with a partially tidy format, the names of the variables are missing form the principal file, and hence we have to extrapolate them from another document. At the end of the process we'll have the following variables:

- Age: age in years
- Sex: (1 = male; 0 = female)
- Chest.Pain: chest pain type Value 1: typical angina Value 2: atypical angina Value 3: non-anginal pain Value 4: Asymptomatic
- BP.Rest: resting blood pressure (in mm Hg on admission to the hospital)
- Chol.Liv: serum cholestoral in mg/dl
- Fast.Blood.Sugar: (fasting blood sugar > 120 mg/dl) (1 = true; 0 = false)
- ECG.Rest: resting electrocardiographic results: Value 0: normal Value 1: having ST-T wave abnormality (T wave inversions and/or ST elevation or depression of > 0.05 mV) Value 2: showing probable or definite left ventricular hypertrophy by Estes' criteria
- HR.Max: maximum heart rate achieved

Loading required package: lattice

- Angina.post.Exercise: exercise induced angina (1 = yes; 0 = no)
- OldPeak.ST: ST depression induced by exercise relative to rest
- $\bullet\,$ Slope. ST: the slope of the peak exercise ST segment
- Vessels.Fluo:number of major vessels (0-3) colored by flourosopy
- Defect:3 = normal; 6 = fixed defect; 7 = reversable defect
- Disease: disease in the patient. It is integer valued from 0 (no presence) to 4. Experiments with the Cleveland database have concentrated on simply attempting to distinguish presence (values 1,2,3,4) from absence (value 0).

```
#Load necessary libraries
if(!require(tidyverse)) install.packages("tidyverse")
## Loading required package: tidyverse
## -- Attaching packages ----
## v ggplot2 3.2.1
                              0.3.2
                     v purrr
## v tibble 2.1.3
                     v dplyr
                              0.8.3
## v tidyr
           0.8.3
                     v stringr 1.4.0
## v readr
           1.3.1
                     v forcats 0.4.0
## -- Conflicts ------
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                   masks stats::lag()
if(!require(caret)) install.packages("caret")
## Loading required package: caret
```

```
##
## Attaching package: 'caret'
## The following object is masked from 'package:purrr':
##
##
       lift
if(!require(data.table)) install.packages("data.table")
## Loading required package: data.table
##
## Attaching package: 'data.table'
## The following objects are masked from 'package:dplyr':
##
##
       between, first, last
## The following object is masked from 'package:purrr':
##
##
       transpose
if(!require(foreign)) install.packages("foreign")
## Loading required package: foreign
if(!require(GGally))install.packages("GGally")
## Loading required package: GGally
## Registered S3 method overwritten by 'GGally':
##
     method from
##
     +.gg
          ggplot2
##
## Attaching package: 'GGally'
## The following object is masked from 'package:dplyr':
##
##
       nasa
#Downloading and checking dataset
data<-read.table(</pre>
"https://archive.ics.uci.edu/ml/machine-learning-databases/heart-disease/processed.cleveland.data",
sep=",")
#adding column names
colnames(data) <- c("Age",</pre>
```

```
"Chest.Pain",
"BP.Rest",
"Chol.Liv",
"Fast.Blood.Sugar",
"ECG.Rest",
"HR.Max",
"Angina.post.Exercise",
"OldPeak.ST",
"Slope.ST",
"Vessels.Fluo",
"Defect",
"Disease")
```

After the first data adjustment we have to check the structure of our datas and check for any NAs (missing values):

```
#Check for changes after renaming columns, and check for NAs str(data)
```

```
## 'data.frame':
                   303 obs. of
                               14 variables:
##
  $ Age
                               63 67 67 37 41 56 62 57 63 53 ...
                         : num
## $ Sex
                               1 1 1 1 0 1 0 0 1 1 ...
                         : num
## $ Chest.Pain
                               1 4 4 3 2 2 4 4 4 4 ...
                         : num
## $ BP.Rest
                               145 160 120 130 130 120 140 120 130 140 ...
                         : num
                               233 286 229 250 204 236 268 354 254 203 ...
## $ Chol.Liv
                         : num
##
  $ Fast.Blood.Sugar
                               1 0 0 0 0 0 0 0 1 ...
                         : num
## $ ECG.Rest
                               2 2 2 0 2 0 2 0 2 2 ...
                         : num
##
   $ HR.Max
                         : num
                               150 108 129 187 172 178 160 163 147 155 ...
## $ Angina.post.Exercise: num 0 1 1 0 0 0 0 1 0 1 ...
                        : num 2.3 1.5 2.6 3.5 1.4 0.8 3.6 0.6 1.4 3.1 ...
## $ OldPeak.ST
## $ Slope.ST
                         : num 3 2 2 3 1 1 3 1 2 3 ...
   $ Vessels.Fluo
                        : Factor w/ 5 levels "?","0.0","1.0",...: 2 5 4 2 2 2 4 2 3 2 ...
## $ Defect
                        : Factor w/ 4 levels "?", "3.0", "6.0", ...: 3 2 4 2 2 2 2 2 4 4 ...
                         : int 0210003021...
##
   $ Disease
#Check for NAs
apply(is.na(data), 2,which)
```

integer(0)

Despite after check there wasn't any NAs, we have seen that some variables was coded by the question mark "?", so few entries are missed. For thi reason we procede to remove them all by filtering dataset.

```
data <- data %>% filter_all(all_vars(.!="?"))
```

The dataset contains the column disease, that is represented by a range of five values 0 to 4, where 0 means "No disease" and the other 4 values are the level of disease progression. Since also a a value of 1 indicates heart disease, we'll convert it to a binary variable where 0 is kept if no disease are present, and 1 otherwise. Additionally, some data loaded as numeric are converted to factor for better analysis.

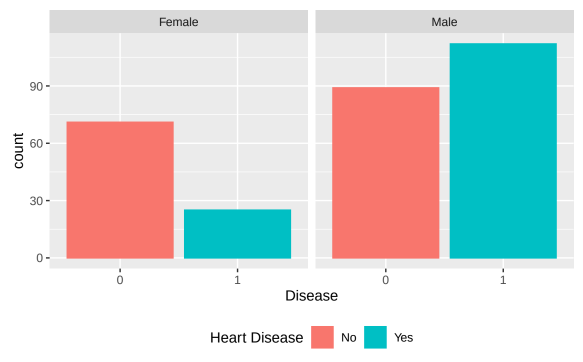
```
data$Vessels.Fluo<-as.numeric(data$Vessels.Fluo) -2
data$Defect<-factor(data$Defect)
data$Sex<-factor(data$Sex)
data$Chest.Pain<-factor(data$Chest.Pain)
data$Fast.Blood.Sugar<-as.factor(data$Fast.Blood.Sugar)
data$ECG.Rest<-as.factor(data$ECG.Rest)
data$Angina.post.Exercise<-as.factor(data$Angina.post.Exercise)
data$Slope.ST<-as.factor(data$Slope.ST)</pre>
data<-data%>%mutate(Disease=as.factor(ifelse(Disease==0,0,1)))
```

Once we have finally cleaned our dataset, we are ready to starting the explorative analysis.

Data Exploration and testing correlations

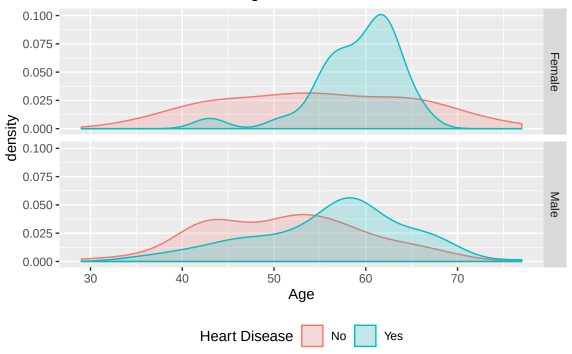
To check our data, will perform a series of density plot and bar plot to better exploring the nature of our independent variables compared to the dependent one (disease). Since epidemiologically it is known that the male and the female have a different distribution of heart diseases, we will plot all the variables comparing male and female sex.

Heart Disease Males VS Females



How is possible to see from the first barplot, males have a bigger incidence and a bigger prevalence of heart disease than females.

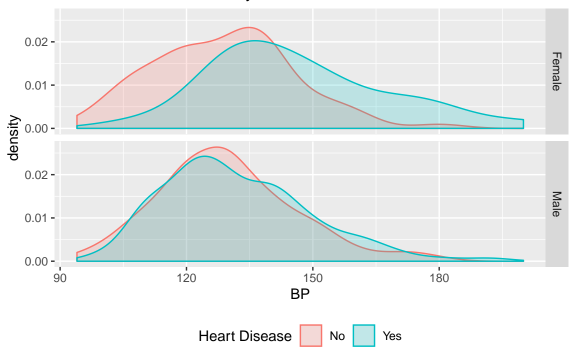
Heart Diseases versus Age for Males and Females



Nο

only the sex represent a discriminant variable for the occurrence of heart disease, age too has an important effect on it. Males and females, as is shown in the previous density plot, are both suffer a greater incidence of heart disease between the age of 50 and 70 years old.

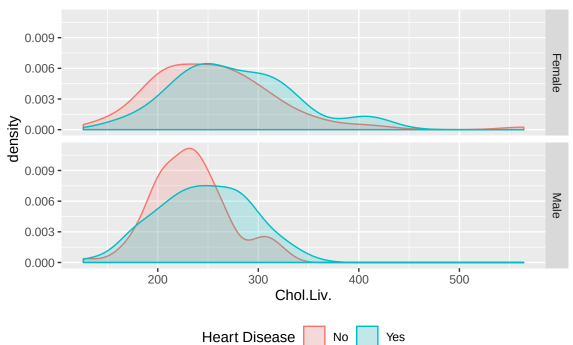
Heart Diseases versus Systolic BP for Males and Females



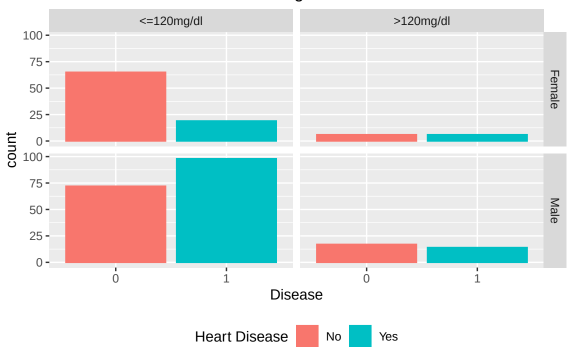
If we

consider the systolic BP we can observe that there aren't great difference in terms of presence of disease, just in case of female sex, we can note a slightly increasing of incidence with a value of 150 mmHg or greater.

Heart Diseases versus Cholesterol Levels for Males and Females



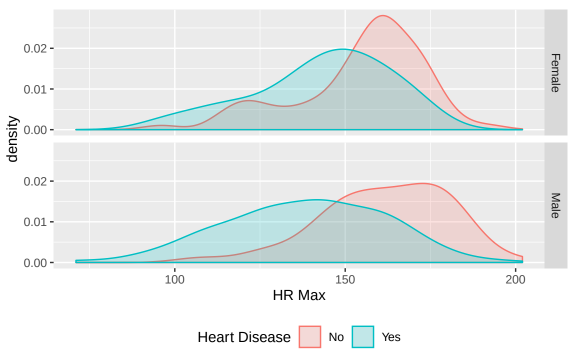
Heart Disease VS FBS > 120 mg/dl Males and Females



Choles-

terol levels and Fast blood sugar are also seeming a non-discriminating variables, although we can observe a little modification in terms of incidence with values of Cholesterol greater than 250.

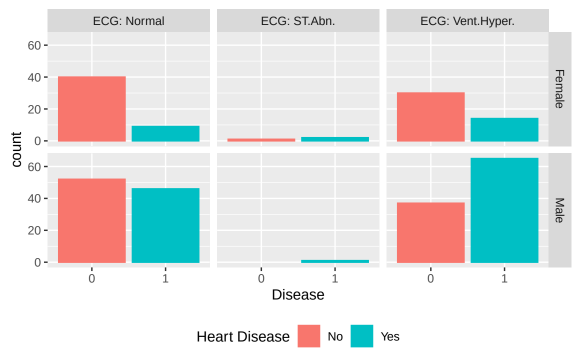
Heart Diseases versus HR max for Males and Females



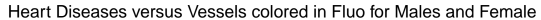
Para-

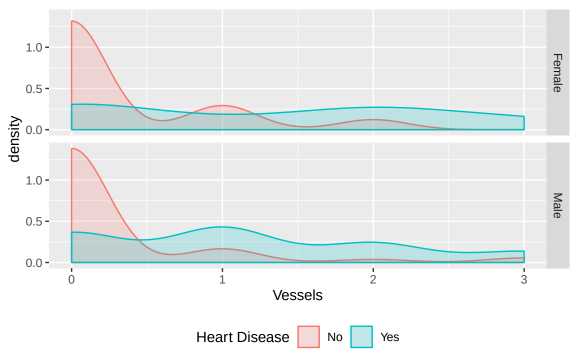
doxically, as is shown in the density plot, a lower heart rate would seem to be related to an increase in the incidence of the disease.

Heart Disease VS ECG Males and Females



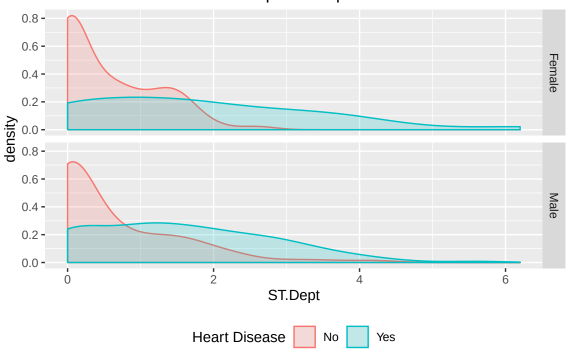
ECG seems to provide insights just for evident anomalies of cardiac rhythm.



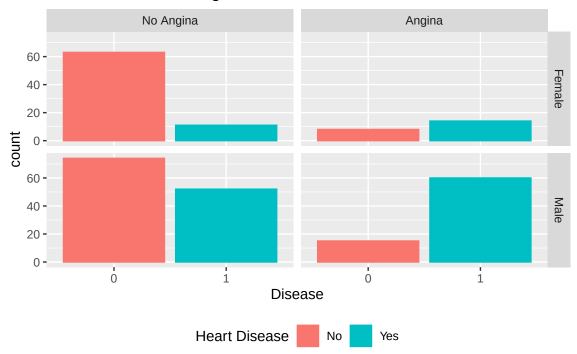


A very important information is provided by the cardiac fluor angiography. In Fact, with one or more vessels involved the incidence of heart disease increase significatively.

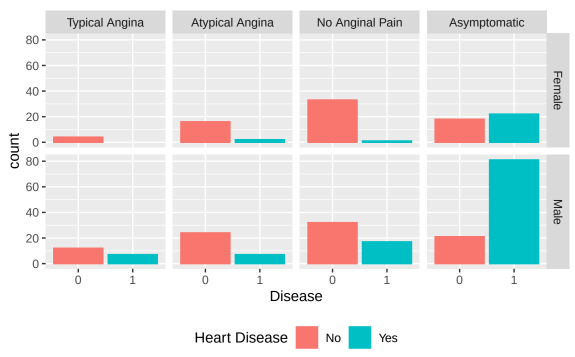




Heart Disease VS Angina Males and Females



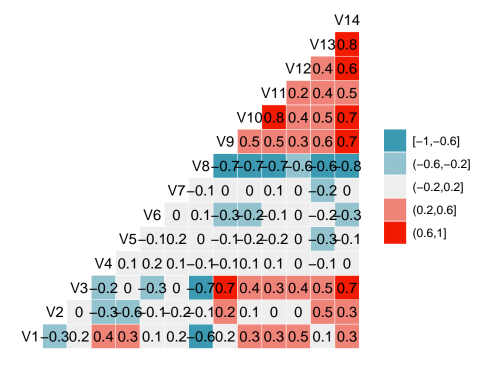
Heart Disease VS Chest Pain Males and Females



While ST depression during an ECG, and an Angina pain after exercise would seem to be related to the presence of heart disease, the data about chest pain shows a counterintuitive condition, where typical angina seems to be a worst predicting element than asymptomatic condition.

Once reached this point of the analysis it is advisable to evaluate the correlations existing between the vari-

Correlation Plot



ables examined.

As we can observe our variables are not very intercorrelated, and so we can proceed with our analysis without the risk of use two or more unreliable predictors.

Creating the Training and Testing Sets

In order to predict heart disease in patients, we must separate the dataset into a training and a testing set, each containing different observations. 20% of the dataset is thus assigned to the testing set.

```
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6

## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used

test_index<-createDataPartition(data$Disease,times = 1, p=0.2, list=F) #Indexing for test set
test_set<-data[test_index,] #creating test set
train_set<-data[-test_index,] #crating training set</pre>
```

We are now ready to test some machine learning algorithms to create a final prediction model for the occurrence of heart diseas, but before start we must check for the integrity of the two sets, and for their homogeneicity.

```
#Checking for datasets distributions
summary(train_set)
## Age Sex Chest.Pain BP.Rest Chol.Liv
## Min. :29.00 0: 78 1: 19 Min. : 94.0 Min. :131.0
```

```
1st Qu.:48.00
                                          1st Qu.:120.0
                                                           1st Qu.:211.0
                     1:159
                              2: 42
##
    Median :56.00
                              3: 65
                                         Median :130.0
                                                           Median :243.0
                                                 :131.2
    Mean
           :54.37
                              4:111
                                          Mean
                                                           Mean
                                                                   :248.1
##
    3rd Qu.:61.00
                                          3rd Qu.:140.0
                                                           3rd Qu.:275.0
##
    Max.
            :74.00
                                         Max.
                                                 :192.0
                                                           Max.
                                                                   :564.0
##
    Fast.Blood.Sugar ECG.Rest
                                                 Angina.post.Exercise
                                    HR.Max
##
    0:206
                      0:116
                                       : 71.0
                                                 0:159
                                Min.
##
    1: 31
                      1: 2
                                1st Qu.:138.0
                                                 1: 78
##
                      2:119
                                Median :153.0
##
                                Mean
                                        :149.8
##
                                3rd Qu.:165.0
##
                                Max.
                                        :202.0
##
      OldPeak.ST
                                Vessels.Fluo
                                                 Defect
                     Slope.ST
                                                            Disease
##
    Min.
            :0.000
                     1:111
                               Min.
                                       :0.0000
                                                 3.0:128
                                                            0:128
##
    1st Qu.:0.000
                     2:112
                               1st Qu.:0.0000
                                                 6.0: 15
                                                            1:109
    Median :0.800
                     3: 14
                               Median :0.0000
                                                 7.0: 94
##
    Mean
            :1.041
                               Mean
                                       :0.6582
                               3rd Qu.:1.0000
    3rd Qu.:1.600
                               Max.
##
    Max.
            :6.200
                                       :3.0000
```

summary(test_set)

```
BP.Rest
                                                             Chol.Liv
##
                     Sex
                             Chest.Pain
         Age
           :37.00
##
    Min.
                     0:18
                             1: 4
                                                                  :126.0
                                         Min.
                                                :100.0
                                                          Min.
    1st Qu.:47.50
                     1:42
                             2: 7
                                         1st Qu.:120.0
                                                          1st Qu.:211.8
    Median :57.00
                             3:18
                                         Median :131.0
                                                          Median :243.5
##
    Mean
            :55.23
                             4:31
                                         Mean
                                                :133.5
                                                          Mean
                                                                  :244.5
##
    3rd Qu.:60.50
                                         3rd Qu.:145.2
                                                          3rd Qu.:282.0
##
    Max.
            :77.00
                                                :200.0
                                                          Max.
                                                                  :327.0
                                    HR.Max
##
    Fast.Blood.Sugar ECG.Rest
                                                 Angina.post.Exercise
##
    0:48
                      0:31
                                Min.
                                        : 95.0
                                                 0:41
##
    1:12
                                1st Qu.:125.0
                                                 1:19
                      1: 2
##
                      2:27
                                Median :152.0
##
                                        :148.8
                                Mean
##
                                3rd Qu.:171.2
##
                                Max.
                                        :194.0
##
      OldPeak.ST
                     Slope.ST
                                Vessels.Fluo
                                               Defect
                                                         Disease
##
                     1:28
                                               3.0:36
                                                         0:32
    Min.
            :0.000
                               Min.
                                       :0.00
##
    1st Qu.:0.000
                     2:25
                               1st Qu.:0.00
                                               6.0: 3
                                                         1:28
##
    Median :0.700
                     3: 7
                               Median:0.00
                                               7.0:21
    Mean
           :1.112
                               Mean
                                       :0.75
##
    3rd Qu.:1.700
                               3rd Qu.:1.00
    Max.
            :5.600
                               Max.
                                       :3.00
```

For a better develop and train our models we will provide to set a training control with the caret package, using a 10 fold cross validation during all over the tests.

```
# Define train control for k-fold (10-fold here) cross validation
fitControl <- trainControl(method = "cv", number = 10, savePredictions = TRUE)</pre>
```

Model 1: K-Nearest Neighbors

A first attempt was made using a k-nearest neighbors algorithm, or 'KNN'. The first step to optimize this model was to use the tuneGrid function in order to retrieve the highest accuracy from a range of k values

(the number of "neighbors" to be considered for each datapoint).

```
#starting with a KNN Model
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6</pre>
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
fit_KNN<-train(Disease~.,</pre>
               data= train_set,
               method="knn",
               tuneGrid=data.frame(k=seq(1,30,1)),
               trControl=fitControl) #fit model and try some k's
pre_KNN<-predict(fit_KNN,test_set) #predict results</pre>
confusionMatrix(pre_KNN,test_set$Disease) #make a confusion matrix
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 23 13
##
            1 9 15
##
##
##
                  Accuracy : 0.6333
                    95% CI: (0.499, 0.7541)
##
##
       No Information Rate: 0.5333
       P-Value [Acc > NIR] : 0.0766
##
##
                     Kappa : 0.2568
##
##
##
   Mcnemar's Test P-Value: 0.5224
##
               Sensitivity: 0.7188
##
##
               Specificity: 0.5357
##
            Pos Pred Value: 0.6389
##
            Neg Pred Value: 0.6250
##
                Prevalence: 0.5333
            Detection Rate: 0.3833
##
##
      Detection Prevalence: 0.6000
##
         Balanced Accuracy: 0.6272
##
##
          'Positive' Class: 0
##
## # A tibble: 1 x 5
##
     Method Accuracy Sensitivity Specificity Bal. Accuracy
##
     <chr>>
               <dbl>
                            <dbl>
                                        <dbl>
                                                      <dbl>
                                        0.536
                                                      0.627
## 1 KNN
               0.633
                            0.719
```

At a first look the accuracy of this model is not very high (0.63) and in particular his Balanced Accuracy is only 0.627, all this things suggest us to continue our analyses.

Model 2: Adaptive boosting Model

The second model that we wanna try is the "Adaboost" model. Also this time we set a seed equal to 1 ande proceed to train the model on the training set. It will take few minutes to run, but crossing our fingers probably we will see an improvement.

```
#Adaboost Classification trees model
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6</pre>
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
fit_ada<-train(Disease~.,</pre>
               data=train_set,
               method="adaboost",
               trControl=fitControl) #fit model
pre_ada<-predict(fit_ada,test_set) #predict results</pre>
confusionMatrix(pre_ada,test_set$Disease) #make a confusion matrix
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 27 7
##
##
            1 5 21
##
##
                  Accuracy: 0.8
                    95% CI : (0.6767, 0.8922)
##
##
       No Information Rate: 0.5333
       P-Value [Acc > NIR] : 1.609e-05
##
##
##
                     Kappa: 0.5964
##
##
    Mcnemar's Test P-Value: 0.7728
##
##
               Sensitivity: 0.8438
##
               Specificity: 0.7500
##
            Pos Pred Value: 0.7941
            Neg Pred Value: 0.8077
##
##
                Prevalence: 0.5333
            Detection Rate: 0.4500
##
##
      Detection Prevalence: 0.5667
##
         Balanced Accuracy: 0.7969
##
##
          'Positive' Class: 0
##
## # A tibble: 2 x 5
##
    Method
              Accuracy Sensitivity Specificity Bal. Accuracy
                                          <dbl>
##
     <chr>>
                 <dbl>
                              <dbl>
                                                        <dbl>
## 1 KNN
                 0.633
                              0.719
                                          0.536
                                                        0.627
## 2 Adaboost
                                          0.75
                 0.8
                              0.844
                                                        0.797
```

We got a good result with an accuracy of 0.8 and a Balanced Accuracy of 0.796 (we note just a slightly lack in the specificity of this model). The results are good, but we want improve it and try to get a better prediction, hence we will train another model on our dataset.

Model 3: General Logistic Regression model

A tibble: 3 x 5

At this point we want to change our strategy and we will try a regression type model, in particular will perform a logistic regression model and will check if it will yield the hoping improvements.

```
#Logistic Regression Model
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6</pre>
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
fit_GLM<-train(Disease~.,</pre>
               data= train_set,
               method="glm",
               family="binomial",
               trControl=fitControl) #fit model
pre_GLM<-predict(fit_GLM,test_set) #predict results</pre>
confusionMatrix(pre_GLM,test_set$Disease) #make a confusion matrix
## Confusion Matrix and Statistics
##
             Reference
##
## Prediction 0 1
##
            0 26 5
##
            1 6 23
##
##
                  Accuracy : 0.8167
                    95% CI: (0.6956, 0.9048)
##
##
       No Information Rate: 0.5333
       P-Value [Acc > NIR] : 4.344e-06
##
##
##
                     Kappa: 0.6325
##
##
    Mcnemar's Test P-Value : 1
##
##
               Sensitivity: 0.8125
##
               Specificity: 0.8214
            Pos Pred Value: 0.8387
##
            Neg Pred Value: 0.7931
##
##
                Prevalence: 0.5333
##
            Detection Rate: 0.4333
##
      Detection Prevalence: 0.5167
         Balanced Accuracy: 0.8170
##
##
##
          'Positive' Class: 0
##
```

##	Method	Accuracy	Sensitivity	Specificity	Bal.Accuracy
##	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
## 1	KNN	0.633	0.719	0.536	0.627
## 2	Adaboost	0.8	0.844	0.75	0.797
## 3	General Logistic Regressio~	0.817	0.812	0.821	0.817

Comparing all the three models done we can see that GLM model improve considerly the values of Accuracy, Sensitivty and Specificity. So, we are on the good way, we could stop our research, but we are "hungry" researcher and we want to check if it is still possible to improve our results.

Model 4: Decision Tree model

##

##

Detection Prevalence: 0.6000 Balanced Accuracy: 0.7946

As fourth model we will choose a decision tree model, and also for this algorithm will use the caret package with a 10 K-Fold cross validation.

```
#Decision tree Model
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6</pre>
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
fit_rpart<-train(Disease~.,</pre>
                 data= train_set,
                 method="rpart",
                 trControl=fitControl) #fit model
pre_rpart<-predict(fit_rpart,test_set) #predict results</pre>
confusionMatrix(pre rpart,test set$Disease) #make a confusion matrix
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 28 8
##
##
            1 4 20
##
##
                  Accuracy: 0.8
                    95% CI : (0.6767, 0.8922)
##
##
       No Information Rate: 0.5333
##
       P-Value [Acc > NIR] : 1.609e-05
##
                     Kappa: 0.5946
##
##
    Mcnemar's Test P-Value: 0.3865
##
##
##
               Sensitivity: 0.8750
##
               Specificity: 0.7143
##
            Pos Pred Value: 0.7778
##
            Neg Pred Value: 0.8333
##
                Prevalence: 0.5333
##
            Detection Rate: 0.4667
```

```
##
##
          'Positive' Class: 0
##
## # A tibble: 4 x 5
     Method
                                   Accuracy Sensitivity Specificity Bal. Accuracy
##
     <chr>
##
                                      <dbl>
                                                   <dbl>
                                                                <dbl>
                                                                              <dbl>
## 1 KNN
                                      0.633
                                                   0.719
                                                                0.536
                                                                             0.627
## 2 Adaboost
                                      0.8
                                                   0.844
                                                                0.75
                                                                             0.797
## 3 General Logistic Regressio~
                                      0.817
                                                   0.812
                                                                0.821
                                                                             0.817
## 4 Decision Tree Model
                                      0.8
                                                   0.875
                                                                0.714
                                                                             0.795
```

Again, checking the results we see a slightly worsen value of accuracy, but despite a lack in specificity we had sensibly improve our prediction sessibility; for this reason, seen the possibility to increase our predictive parameters, it has meant continuing our analysis trying to improve the specificity as well as the sensitivity.

Model 5: Random Forest model

##

As further model we will train a Random Forest model, that represents a deepening of the Decision Tree model seen before.

```
#Random Forest model
set.seed(1, sample.kind = "Rounding") #use set.seed(1) if use R version < 3.6</pre>
## Warning in set.seed(1, sample.kind = "Rounding"): non-uniform 'Rounding'
## sampler used
fit_rf<-train(Disease~.,</pre>
              data= train_set,
              method="rf",
              trControl=fitControl) #fit model
pre_rf<-predict(fit_rf,test_set) #predict results</pre>
confusionMatrix(pre_rf,test_set$Disease) #make a confusion matrix
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction
              0 1
            0 28 5
##
            1 4 23
##
##
                  Accuracy: 0.85
##
                    95% CI : (0.7343, 0.929)
##
       No Information Rate: 0.5333
##
       P-Value [Acc > NIR] : 2.293e-07
##
##
##
                      Kappa: 0.698
##
   Mcnemar's Test P-Value : 1
##
##
```

Sensitivity: 0.8750

```
##
               Specificity: 0.8214
##
            Pos Pred Value: 0.8485
##
            Neg Pred Value: 0.8519
                Prevalence: 0.5333
##
##
            Detection Rate: 0.4667
##
      Detection Prevalence: 0.5500
         Balanced Accuracy: 0.8482
##
##
##
          'Positive' Class: 0
##
## # A tibble: 5 x 5
     Method
                                  Accuracy Sensitivity Specificity Bal. Accuracy
##
##
     <chr>>
                                     <dbl>
                                                  <dbl>
                                                               <dbl>
                                                                            <dbl>
## 1 KNN
                                     0.633
                                                  0.719
                                                               0.536
                                                                            0.627
## 2 Adaboost
                                     0.8
                                                  0.844
                                                              0.75
                                                                            0.797
## 3 General Logistic Regressio~
                                     0.817
                                                  0.812
                                                              0.821
                                                                            0.817
## 4 Decision Tree Model
                                     0.8
                                                  0.875
                                                               0.714
                                                                            0.795
## 5 Random Forest
                                     0.85
                                                  0.875
                                                               0.821
                                                                            0.848
```

Our efforts are payed with a strong increase of accuracy (0.85), sensitivty (0.875) and specificity (0.821). All our parmeters are improved and now, once reached the decision point, we have to choose if stop the searching, or try to make some changes to improve again. We decide to try another strategy, will ensemble the three best models to see if is possible to get a more accurate result.

Ensemble model

The ensemble strategy involves calculating the average of the results obtained with the desired models, to generate a new prediction series, that will be subsequently compared with the real results through a confusion matrix.

```
#Try to improve our models with ensemble strategy
ensemble<-data.frame(pre_GLM,pre_rf,pre_ada)
pre_ensemble<-as.factor(ifelse(rowMeans(ensemble==1)>0.5,1,0))
confusionMatrix(pre_ensemble,test_set$Disease)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0 1
            0 28 5
##
            1 4 23
##
##
##
                  Accuracy: 0.85
##
                    95% CI: (0.7343, 0.929)
##
       No Information Rate: 0.5333
##
       P-Value [Acc > NIR] : 2.293e-07
##
##
                     Kappa: 0.698
##
    Mcnemar's Test P-Value : 1
##
##
```

```
##
               Sensitivity: 0.8750
##
               Specificity: 0.8214
            Pos Pred Value: 0.8485
##
##
            Neg Pred Value: 0.8519
##
                Prevalence: 0.5333
            Detection Rate: 0.4667
##
##
      Detection Prevalence: 0.5500
         Balanced Accuracy: 0.8482
##
##
##
          'Positive' Class : 0
##
## # A tibble: 6 x 5
                                  Accuracy Sensitivity Specificity Bal. Accuracy
##
     Method
##
     <chr>>
                                      <dbl>
                                                  <dbl>
                                                               <dbl>
                                                                            <dbl>
## 1 KNN
                                     0.633
                                                  0.719
                                                               0.536
                                                                            0.627
## 2 Adaboost
                                     0.8
                                                  0.844
                                                               0.75
                                                                            0.797
## 3 General Logistic Regressio~
                                     0.817
                                                  0.812
                                                               0.821
                                                                            0.817
## 4 Decision Tree Model
                                     0.8
                                                  0.875
                                                               0.714
                                                                            0.795
## 5 Random Forest
                                     0.85
                                                  0.875
                                                               0.821
                                                                            0.848
## 6 Ensamble Model
                                     0.85
                                                  0.875
                                                               0.821
                                                                            0.848
```

Conclusions

In conclusion we can say that we have achieved good predictive results through the Random Forest model. The levels of accuracy, sensitivity and specificity are quite satisfactory and allow to predict in 85% of cases the onset of heart disease in both men and women. Probably further variables to be examined could positively influence forecasts and further improve our machine learning models.

[&]quot;' As we can see the ensemble model didn't improve our results achieved from the Random Forest, so we can stop to search another algorithm or another strategy and choose as definitive approach the Random Forest model.