CS5525 Final Project Code Submission

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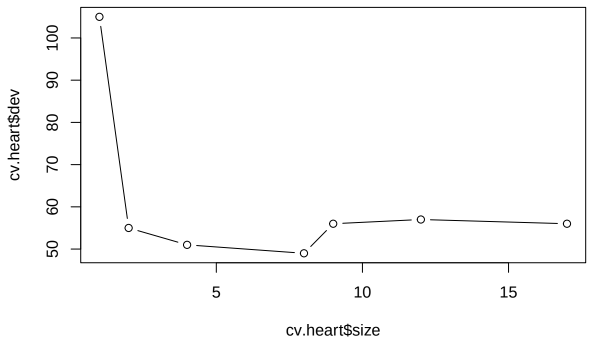
Classification Methods

First set the directory (path which contains the heart.csv data), and import any needed libraries

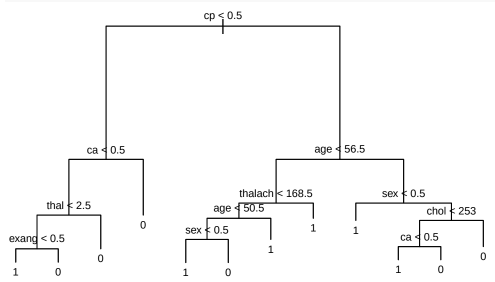
```
#setwd(" ") # uncomment to set working directory via code
library(tree)
library(randomForest) # bootsrap/bagging & random forest
library(class) # KNN
library(caret) # SVM
library(ggplot2)
library(cowplot)
```

Decision Trees

```
## Classification tree:
## tree(formula = Target ~ . - target, data = heart, subset = train)
## Variables actually used in tree construction:
## [1] "cp"
                 "ca"
                           "thal"
                                     "exang"
                                                "age"
                                                          "oldpeak" "chol"
## [8] "thalach" "sex"
## Number of terminal nodes: 17
## Residual mean deviance: 0.4429 = 93 / 210
## Misclassification error rate: 0.1145 = 26 / 227
## -- Plot tree
plot(tree.heart)
text(tree.heart, pretty=1, cex=0.7)
                                                 ca < 0.5
                                       thalach < 168.5
                                                         sex ₹ 0.5
                    oldpeak < 0.45
                                     age ₹ 50.5
                                                   age < 62
                                                               chol k 253
                         sex < 0.5
age < 60
                                       0
             0
                 0
# Prune the classification tree
set.seed(2441139)
cv.heart <- cv.tree(tree.heart, FUN=prune.misclass)</pre>
cv.heart
## $size
## [1] 17 12 9 8 4 2 1
##
## $dev
## [1] 56 57 56 49 51 55 105
##
## $k
            -Inf 0.0000000 0.6666667 1.0000000 3.0000000 7.0000000 50.00000000
## [1]
## $method
## [1] "misclass"
##
## attr(,"class")
## [1] "prune"
                       "tree.sequence"
plot(cv.heart$size, cv.heart$dev, type='b')
```



```
prune.heart <- prune.misclass(tree.heart, best=10)
plot(prune.heart)
text(prune.heart, pretty=1, cex=0.65)</pre>
```

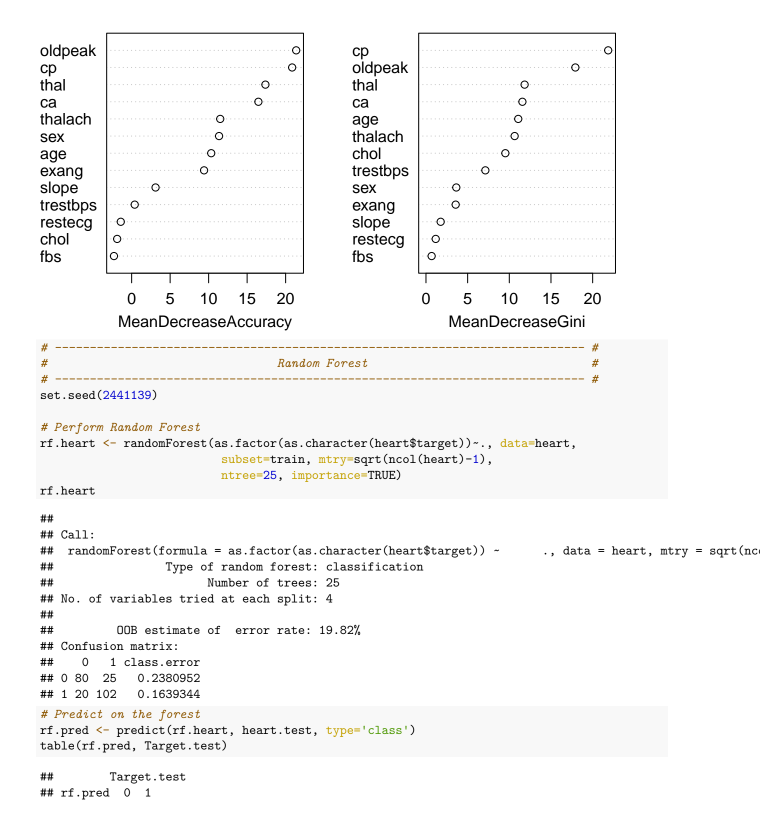


```
# Predict using test set and pruned tree. Compare.
tree.pred <- predict(tree.heart, heart.test, type='class')  # test tree
prune.pred <- predict(prune.heart, heart.test, type='class')  # pruned tree
table(prune.pred, Target.test)</pre>
```

```
## Target.test
## prune.pred 0 1
## 0 28 11
## 1 5 32
```

```
table(tree.pred, Target.test)
           Target.test
## tree.pred 0 1
          0 28 11
##
##
          1 5 32
                                  Bagging
set.seed(2441139)
# Perform bagging
bag.heart <- randomForest(as.factor(as.character(heart$target))~., data=heart,</pre>
                         subset=train, mtry=ncol(heart)-1,
                          importance=TRUE)
bag.heart
##
## Call:
## randomForest(formula = as.factor(as.character(heart$target)) ~ ., data = heart, mtry = ncol(he
##
                 Type of random forest: classification
                       Number of trees: 500
## No. of variables tried at each split: 13
##
          OOB estimate of error rate: 22.91%
##
## Confusion matrix:
## 0 1 class.error
## 0 75 30 0.2857143
## 1 22 100 0.1803279
# Predict on bagged tree
bag.pred <- predict(bag.heart, heart.test, type='class')</pre>
table(bag.pred, Target.test)
          Target.test
##
## bag.pred 0 1
##
         0 27 7
         1 6 36
##
varImpPlot(bag.heart)
```

bag.heart



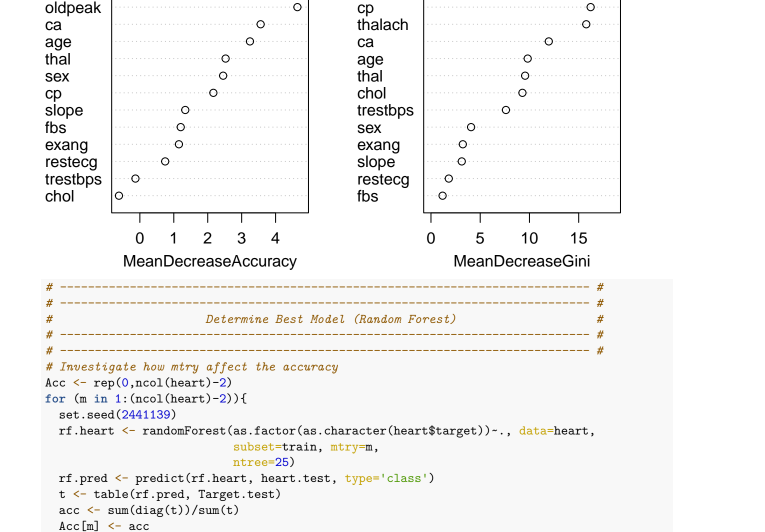
```
## 0 30 6
## 1 3 37
varImpPlot(rf.heart)
```

oldpeak

rf.heart

thalach

mbest <- which(Acc==max(Acc))</pre>

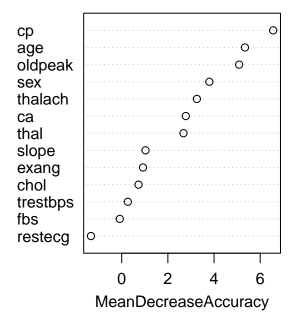


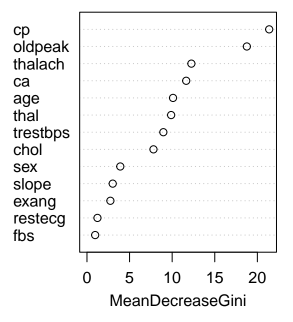
plot(1:(ncol(heart)-2), Acc, xlab='mtry', ylab='Accuracy of random forest') # include plot in final sub

```
0.90
                                                                                   0
Accuracy of random forest
                                                                                                                                       0
         0.85
                                                                                             0
                     0
                               0
                                                    0
                                                                         0
                                                                                                                            0
         0.80
                                          0
                                                                                                        0
         0.75
                                                                                                                  0
                               2
                                                    4
                                                                         6
                                                                                             8
                                                                                                                 10
                                                                                                                                      12
                                                                           mtry
```

```
# Now use the best value of m for the random forest
set.seed(2441139)
rf.heart <- randomForest(as.factor(as.character(heart$target))~., data=heart,
                         subset=train, mtry=mbest,
                         ntree=25, importance=TRUE)
rf.heart
##
## Call:
##
    randomForest(formula = as.factor(as.character(heart$target)) ~ ., data = heart, mtry = mbest,
##
                  Type of random forest: classification
                        Number of trees: 25
## No. of variables tried at each split: 7
##
           OOB estimate of error rate: 23.79%
##
## Confusion matrix:
##
      0 1 class.error
## 0 78 27
             0.2571429
## 1 27 95
            0.2213115
# Predict on the forest
rf.pred <- predict(rf.heart, heart.test, type='class')</pre>
table(rf.pred, Target.test)
##
          Target.test
## rf.pred 0 1
##
         0 28 7
         1 5 36
varImpPlot(rf.heart)
```

rf.heart





KNN

```
# K-Nearest Neighbor
cl <- as.factor(heart$target[train])
knn.heart <- knn(heart[train,], heart.test, cl, k = 5, prob=TRUE)
table(knn.heart, Target.test)

## Target.test
## knn.heart 0 1
## 0 17 15
## 1 16 28</pre>
```

SVM

```
set.seed(2441139)

# splitting data into test and train
intrain <- createDataPartition(y = heart$target, p = 0.7, list = F)

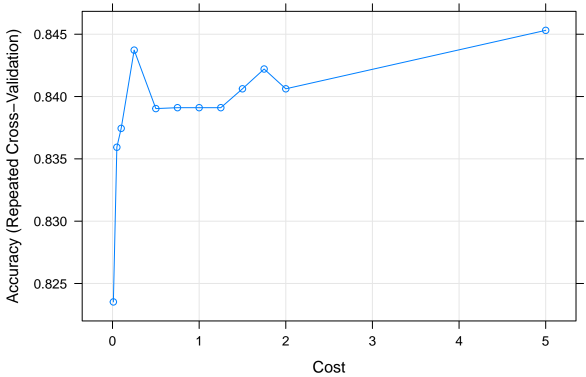
training <- heart[intrain,]
testing <- heart[-intrain,]
training[["target"]] <- as.factor(training[["target"]])

dim(training)</pre>
```

[1] 213 14

```
dim(testing)
## [1] 90 14
# model training with sum
trctrl <- trainControl(method = "repeatedcv", number = 10, repeats = 3)</pre>
svm.mod <- train(target ~ ., data = training, method = "svmLinear",</pre>
                trControl = trctrl,
                preProcess = c("center", "scale"),
                tuneLength = 10)
svm.mod
## Support Vector Machines with Linear Kernel
##
## 213 samples
## 13 predictor
   2 classes: '0', '1'
##
## Pre-processing: centered (13), scaled (13)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 192, 191, 192, 192, 191, 191, ...
## Resampling results:
##
##
    Accuracy
               Kappa
    0.8266955 0.6511675
##
##
## Tuning parameter 'C' was held constant at a value of 1
# predction using the above model
svm.pred <- predict(svm.mod, newdata = testing)</pre>
svm.pred
## [39] 1 1 1 1 1 1 1 0 0 1 1 1 1 1 0 1 0 1 1 1 1 0 0 0 0 0 0 0 0 0 0 0 1 0 0 0 0 0 0
## [77] 0 0 0 1 1 1 0 0 1 0 1 0 0 0
## Levels: 0 1
# accuracy of the trained model
confusionMatrix(table(svm.pred, testing$target))
## Confusion Matrix and Statistics
##
##
## svm.pred 0 1
         0 28 11
##
##
         1 9 42
##
##
                 Accuracy : 0.7778
##
                   95% CI: (0.6779, 0.8587)
##
      No Information Rate: 0.5889
##
      P-Value [Acc > NIR] : 0.0001266
##
##
                    Kappa: 0.5448
##
##
   Mcnemar's Test P-Value: 0.8230633
##
```

```
##
               Sensitivity: 0.7568
##
               Specificity: 0.7925
##
            Pos Pred Value: 0.7179
##
            Neg Pred Value: 0.8235
##
                Prevalence: 0.4111
##
            Detection Rate: 0.3111
##
      Detection Prevalence: 0.4333
##
         Balanced Accuracy: 0.7746
##
##
          'Positive' Class: 0
##
# costs for further tuning with 10-fold cross-validation
grid \leftarrow expand.grid(C = c(0, 0.01, 0.05, 0.1, 0.25, 0.5, 0.75, 1, 1.25, 1.5, 1.75, 2, 5))
svm.mod.grid <- train(target ~ ., data = training, method = "svmLinear",</pre>
                      trControl = trctrl,
                      preProcess = c("center", "scale"),
                      tuneGrid = grid,
                      tuneLength = 10)
svm.mod.grid
## Support Vector Machines with Linear Kernel
##
## 213 samples
  13 predictor
##
    2 classes: '0', '1'
##
## Pre-processing: centered (13), scaled (13)
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 192, 191, 192, 192, 192, 192, ...
## Resampling results across tuning parameters:
##
##
     C
           Accuracy
##
    0.00
                 {\tt NaN}
                            NaN
##
    0.01 0.8235209 0.6414705
##
    0.05 0.8359307 0.6677455
          0.8374459 0.6712576
##
     0.10
##
     0.25 0.8437229 0.6843572
##
     0.50 0.8390332 0.6749413
     0.75 0.8391053 0.6752263
##
##
     1.00 0.8391053 0.6753687
##
     1.25 0.8391053 0.6753687
##
     1.50 0.8406205 0.6783406
     1.75 0.8422078 0.6815951
##
##
     2.00
          0.8406205
                     0.6785395
##
     5.00 0.8453102 0.6879521
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was C = 5.
# accuracy plot of tuned model
plot(svm.mod.grid)
```



```
# prediction using tuned model
svm.pred.grid <- predict(svm.mod.grid, newdata = testing)</pre>
svm.pred.grid
   ## [39] 1 1 1 1 1 1 1 0 0 1 1 1 1 0 1 0 1 1 1 1 0 0 0 0 0 0 0 0 0 0 0 1 0 0 0 0 0 0
## [77] 0 0 0 1 1 1 0 0 1 0 1 0 0 0
## Levels: 0 1
# accuracy of the tuned model
confusionMatrix(table(svm.pred.grid, testing$target))
## Confusion Matrix and Statistics
##
##
## svm.pred.grid 0 1
##
             0 28 10
##
             1 9 43
##
                Accuracy : 0.7889
##
                  95% CI: (0.6901, 0.8679)
##
##
      No Information Rate: 0.5889
##
      P-Value [Acc > NIR] : 4.918e-05
##
##
                   Kappa: 0.5658
##
##
   Mcnemar's Test P-Value : 1
##
##
             Sensitivity: 0.7568
             Specificity: 0.8113
##
```

Pos Pred Value: 0.7368

##

```
## Neg Pred Value : 0.8269
## Prevalence : 0.4111
## Detection Rate : 0.3111
## Detection Prevalence : 0.4222
## Balanced Accuracy : 0.7840
##
## 'Positive' Class : 0
##
```

Logistic Regression

```
set.seed(2441139)
# pre-processing of data
str(heart)
## 'data.frame':
                    303 obs. of 14 variables:
## $ age : int 63 37 41 56 57 57 56 44 52 57 ...
              : int 1 1 0 1 0 1 0 1 1 1 ...
## $ sex
## $ ср
              : int 3 2 1 1 0 0 1 1 2 2 ...
## $ trestbps: int 145 130 130 120 120 140 140 120 172 150 ...
           : int 233 250 204 236 354 192 294 263 199 168 ...
## $ chol
              : int 100000010...
## $ fbs
## $ restecg : int 0 1 0 1 1 1 0 1 1 1 ...
## $ thalach : int 150 187 172 178 163 148 153 173 162 174 ...
## $ exang : int 000010000...
## $ oldpeak : num 2.3 3.5 1.4 0.8 0.6 0.4 1.3 0 0.5 1.6 ...
## $ slope : int 0 0 2 2 2 1 1 2 2 2 ...
            : int 0000000000...
## $ ca
## $ thal
             : int 1 2 2 2 2 1 2 3 3 2 ...
## $ target : int 1 1 1 1 1 1 1 1 1 ...
heart$sex <- ifelse(test = heart$sex == 0, yes = "F", no = "M")
heart$sex <- as.factor(heart$sex)</pre>
heart$cp <- as.factor(heart$cp)</pre>
heart$fbs <- as.factor(heart$fbs)</pre>
heart$restecg <- as.factor(heart$restecg)</pre>
heart$exang <- as.factor(heart$exang)</pre>
heart$slope <- as.factor(heart$slope)</pre>
heart$ca <- as.factor(heart$ca)</pre>
heart$thal <- as.factor(heart$thal)</pre>
heart$age <- as.numeric(heart$age)</pre>
heart$trestbps <- as.numeric(heart$trestbps)</pre>
heart$chol <- as.numeric(heart$chol)</pre>
heart$thalach <- as.numeric(heart$thalach)</pre>
heart$target <- ifelse(test = heart$target == 0, yes = "Healthy", no = "Unhealthy")
heart$target <- as.factor(heart$target)</pre>
str(heart)
## 'data.frame':
                   303 obs. of 14 variables:
## $ age : num 63 37 41 56 57 57 56 44 52 57 ...
## $ sex
              : Factor w/ 2 levels "F", "M": 2 2 1 2 1 2 1 2 2 2 ...
## $ ср
              : Factor w/ 4 levels "0","1","2","3": 4 3 2 2 1 1 2 2 3 3 ...
## $ trestbps: num 145 130 130 120 120 140 140 120 172 150 ...
```

```
: num 233 250 204 236 354 192 294 263 199 168 ...
## $ fbs
            : Factor w/ 2 levels "0","1": 2 1 1 1 1 1 1 2 1 ...
## $ restecg : Factor w/ 3 levels "0","1","2": 1 2 1 2 2 2 1 2 2 2 ...
## $ thalach : num 150 187 172 178 163 148 153 173 162 174 ...
            : Factor w/ 2 levels "0","1": 1 1 1 1 2 1 1 1 1 1 ...
## $ oldpeak : num 2.3 3.5 1.4 0.8 0.6 0.4 1.3 0 0.5 1.6 ...
## $ slope : Factor w/ 3 levels "0","1","2": 1 1 3 3 3 2 2 3 3 3 ...
             : Factor w/ 5 levels "0","1","2","3",..: 1 1 1 1 1 1 1 1 1 1 ...
## $ thal
             : Factor w/ 4 levels "0","1","2","3": 2 3 3 3 3 2 3 4 4 3 ...
## $ target : Factor w/ 2 levels "Healthy", "Unhealthy": 2 2 2 2 2 2 2 2 2 ...
# getting the number of samples based on gender
xtabs(~ target + sex, data = heart)
##
             sex
                F
## target
##
               24 114
    Healthy
    Unhealthy 72 93
# simple logistic model
logistic <- glm(target ~ sex, data = heart, family = "binomial")</pre>
summary(logistic)
##
## Call:
## glm(formula = target ~ sex, family = "binomial", data = heart)
## Deviance Residuals:
      Min
                1Q
                    Median
                                  3Q
                                          Max
## -1.6651 -1.0923
                    0.7585 1.2650
                                       1.2650
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) 1.0986
                          0.2357 4.661 3.15e-06 ***
                           0.2740 -4.752 2.01e-06 ***
## sexM
               -1.3022
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 417.64 on 302 degrees of freedom
##
## Residual deviance: 392.80 on 301 degrees of freedom
## AIC: 396.8
##
## Number of Fisher Scoring iterations: 4
R_sq_1 <- 1 - logistic$deviance / logistic$null.deviance</pre>
R_sq_1
## [1] 0.05947945
BIC_1 <- logistic$deviance + 2 * log(dim(heart)[1])
BIC_1
## [1] 404.2246
```

```
# complex logistic model
logistic <- glm(target ~ ., data = heart, family = "binomial")</pre>
summary(logistic)
##
## Call:
## glm(formula = target ~ ., family = "binomial", data = heart)
##
## Deviance Residuals:
##
      Min
                               3Q
               1Q
                   Median
                                      Max
## -2.9459 -0.2738
                   0.1012
                          0.4515
                                    3.1248
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) 0.179045 3.705420 0.048 0.961461
                        0.025428 1.094 0.273938
## age
             0.027819
## sexM
             -1.862297
                        0.570844 -3.262 0.001105 **
## cp1
              0.864708
                      0.578000 1.496 0.134645
## cp2
             ## cp3
             2.417107
                        0.719242 3.361 0.000778 ***
             ## trestbps
             -0.004291
## chol
                        0.004245 -1.011 0.312053
## fbs1
             0.445666 0.587977 0.758 0.448472
## restecg1
             0.460582 0.399615 1.153 0.249089
## restecg2
            -0.714204
                        2.768873 -0.258 0.796453
## thalach
             0.020055 0.011859
                                 1.691 0.090820
## exang1
             -0.779111
                        0.451839 -1.724 0.084652 .
## oldpeak
             ## slope1
             -0.775084
                        0.880495 -0.880 0.378707
## slope2
             0.689965 0.947657
                                0.728 0.466568
## ca1
             -2.342301
                        0.527416 -4.441 8.95e-06 ***
## ca2
             -3.483178
                        0.811640 -4.292 1.77e-05 ***
## ca3
             -2.247144 0.937629 -2.397 0.016547 *
## ca4
             1.267961
                        1.720014 0.737 0.461013
## thal1
             2.637558 2.684285 0.983 0.325808
## thal2
             2.367747
                        2.596159 0.912 0.361759
                        2.600380 0.352 0.724901
## thal3
             0.915115
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 417.64 on 302 degrees of freedom
## Residual deviance: 179.63 on 280 degrees of freedom
## AIC: 225.63
##
## Number of Fisher Scoring iterations: 6
R_sq_2 <- 1 - logistic$deviance / logistic$null.deviance</pre>
R_sq_2
## [1] 0.569889
BIC_2 <- logistic$deviance + 14 * log(dim(heart)[1])
BIC_2
```

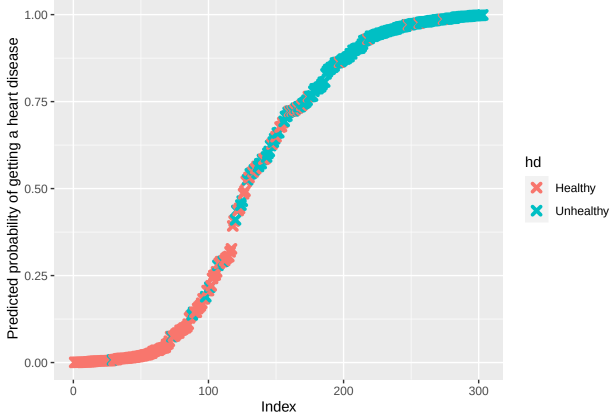
```
## [1] 259.623
```

```
# why age isn't of statistical significance
median(heart$age)
```

[1] 55

```
# plotting the probability of getting a heart disease
predict.hd <- data.frame(prob.of.hd = logistic$fitted.values, hd = heart$target)
predict.hd <- predict.hd[order(predict.hd$prob.of.hd, decreasing = FALSE), ]
predict.hd$rank <- 1:nrow(predict.hd)

ggplot(data = predict.hd, aes(x = rank, y = prob.of.hd)) +
    geom_point(aes(color = hd), alpha = 1, shape = 4, stroke = 2) +
    xlab("Index") +
    ylab("Predicted probability of getting a heart disease")</pre>
```



```
# predicting model accuracy
log.mod <- glm(target ~ ., data = training, family = "binomial")
log.pred <- predict(log.mod, newdata = testing, type = "response")
log.pred <- ifelse(log.pred > 0.5, 1, 0)
confusionMatrix(table(log.pred, testing$target))
```

```
## Confusion Matrix and Statistics
##
##
## log.pred 0 1
## 0 28 11
## 1 9 42
```

Accuracy : 0.7778 95% CI: (0.6779, 0.8587) ## ## No Information Rate : 0.5889 P-Value [Acc > NIR] : 0.0001266 ## ## Kappa : 0.5448 ## ## Mcnemar's Test P-Value : 0.8230633 ## ## Sensitivity: 0.7568 ## ## Specificity: 0.7925 Pos Pred Value : 0.7179 ## Neg Pred Value: 0.8235 ## ## Prevalence : 0.4111 Detection Rate: 0.3111 ## ## Detection Prevalence : 0.4333 ## Balanced Accuracy : 0.7746 ## 'Positive' Class : 0

##