CS 221 Project Final R Code

This R-markdown file contains the R-code for the project report of my CS 221 project.

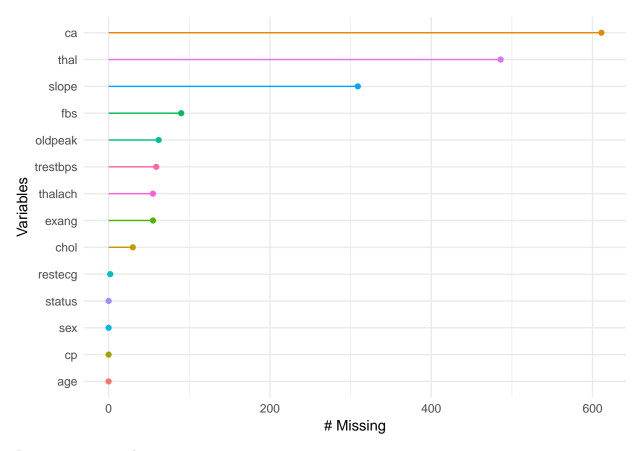
```
library(caTools)
library(caret)
## Loading required package: lattice
## Loading required package: ggplot2
library(e1071)
library(MASS)
library(dplyr)
##
## Attaching package: 'dplyr'
  The following object is masked from 'package:MASS':
##
##
       select
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(ggplot2)
library(naniar)
#library(randomForest)
```

This "convert" helper function is due to Brigitte Mueller, Predicting Heart Disease UCI. I'm borrowing this helper here to change the data type of the Cleveland data read from the website.

The heart disease data are directly read from the UCI website.

```
chclass = c("numeric", "factor", "factor", "numeric", "numeric", "factor", "factor", "numeric", "factor", "factor",
combined.data = convert(combined.data,chclass)
levels(combined.data$status) = c("healthy", "disease")
levels(combined.data$sex) = c("female", "male")
summary(combined.data)
##
                                                                                      trestbps
                                                                                                                          chol
                 age
                                               sex
                                                                 ср
##
     Min.
                    :28.00
                                       female:194
                                                                 1: 46
                                                                                              : 0.0
                                                                                                                Min.
                                                                                                                              : 0.0
                                      male :726
      1st Qu.:47.00
                                                                 2:174
                                                                                1st Qu.:120.0
                                                                                                                1st Qu.:175.0
## Median :54.00
                                                                 3:204
                                                                                Median :130.0
                                                                                                                Median :223.0
                                                                                              :132.1
## Mean
                     :53.51
                                                                 4:496
                                                                                Mean
                                                                                                                Mean
                                                                                                                              :199.1
       3rd Qu.:60.00
                                                                                3rd Qu.:140.0
                                                                                                                3rd Qu.:268.0
                     :77.00
                                                                                              :200.0
                                                                                                                Max.
##
     Max.
                                                                                Max.
                                                                                                                              :603.0
##
                                                                                NA's
                                                                                              :59
                                                                                                                NA's
                                                                                                                              :30
                                                                                                              oldpeak
##
                                                         thalach
           fbs
                             restecg
                                                                                    exang
##
               :692
                                     :551
                                                  Min.
                                                                : 60.0
                                                                                  0
                                                                                           :528
                                                                                                        Min.
                                                                                                                      :-2.6000
##
               :138
                                     :179
                                                   1st Qu.:120.0
                                                                                           :337
                                                                                                        1st Qu.: 0.0000
       1
                             1
                                                                                  1
       NA's: 90
                                     :188
                                                   Median :140.0
                                                                                                        Median : 0.5000
##
                             2
                                                                                  NA's: 55
##
                             NA's: 2
                                                  Mean
                                                                :137.5
                                                                                                        Mean
                                                                                                                    : 0.8788
                                                   3rd Qu.:157.0
                                                                                                        3rd Qu.: 1.5000
##
##
                                                   Max.
                                                                 :202.0
                                                                                                        Max.
                                                                                                                      : 6.2000
##
                                                   NA's
                                                                 :55
                                                                                                        NA's
                                                                                                                      :62
##
                                                       thal
         slope
                                   ca
                                                                                status
##
       1
               :203
                                     :181
                                                   3
                                                           :196
                                                                        healthy:411
                             0
                                                                         disease:509
##
       2
               :345
                             1
                                     : 67
                                                   6
                                                           : 46
                                     : 41
##
       3
               : 63
                                                  7
                                                           :192
                             2
##
      NA's:309
                             3
                                     : 20
                                                  NA's:486
##
                             NA's:611
##
##
str(combined.data)
## 'data.frame':
                                       920 obs. of 14 variables:
                           : num 28 29 29 29 30 31 31 32 32 32 ...
       $ age
                            : Factor w/ 2 levels "female", "male": 2 2 2 2 1 1 2 1 2 2 ...
##
       $ sex
##
       $ cp
                           : Factor w/ 4 levels "1","2","3","4": 2 2 2 2 1 2 4 2 1 2 ...
       $ trestbps: num 130 120 130 140 170 100 120 105 95 110 ...
       $ chol
                           : num 132 243 204 NA 237 219 270 198 0 225 ...
                            : Factor w/ 2 levels "0", "1": 1 1 1 1 1 1 1 1 NA 1 ...
## $ fbs
       $ restecg : Factor w/ 3 levels "0","1","2": 3 1 3 1 2 2 1 1 1 1 ...
## $ thalach : num 185 160 202 170 170 150 153 165 127 184 ...
                         : Factor w/ 2 levels "0", "1": 1 1 1 1 1 2 1 1 1 ...
##
       $ exang
       $ oldpeak : num  0 0 0 0 0 0 1.5 0 0.7 0 ...
                           : Factor w/ 3 levels "1", "2", "3": NA NA 1 NA NA NA 2 NA 1 NA ...
##
       $ slope
## $ ca
                            : Factor w/ 4 levels "0","1","2","3": NA NA 1 NA NA NA NA NA NA NA NA NA ...
                            : Factor w/ 3 levels "3", "6", "7": NA NA 1 NA 2 NA NA NA NA NA ...
## $ thal
       $ status : Factor w/ 2 levels "healthy", "disease": 1 1 1 1 1 1 2 1 2 1 ...
Graphing missing data
```

gg miss var(combined.data)

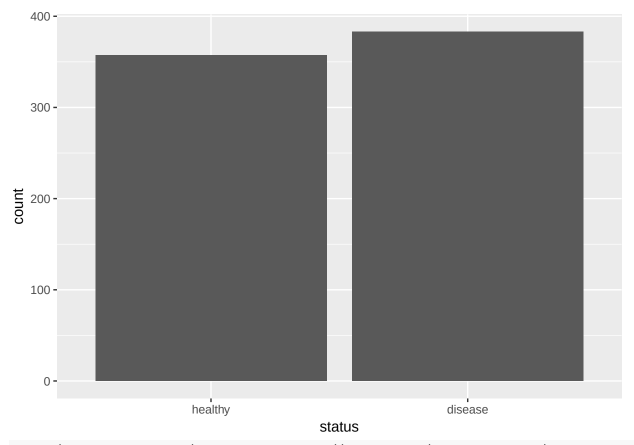


Processing missing data

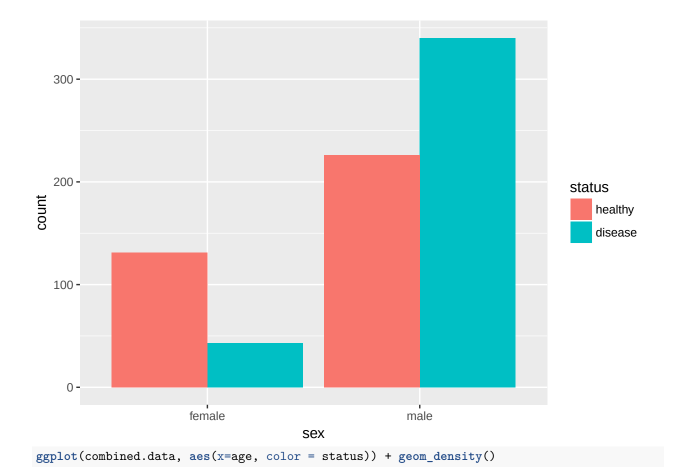
```
combined.data = subset(combined.data, select = -c(11, 12, 13))
combined.data = na.omit(combined.data)
```

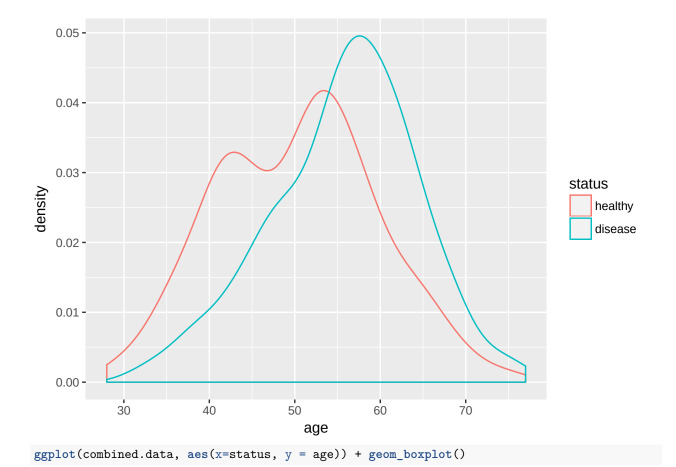
Graphics for data exploration

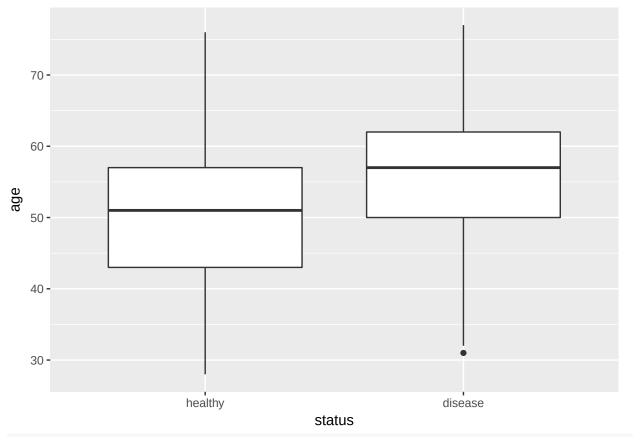
```
ggplot(combined.data, aes(x = status)) + geom_bar()
```



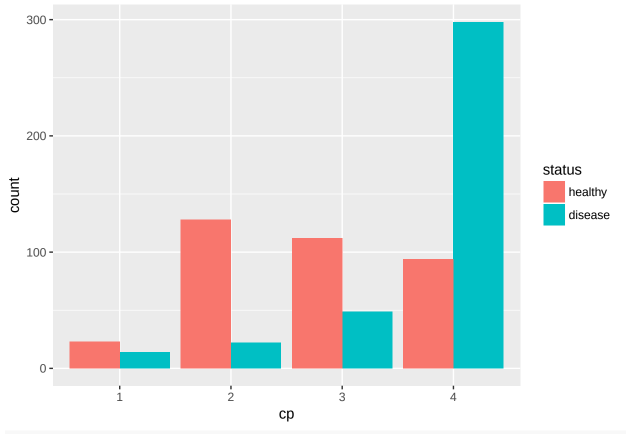
ggplot(combined.data, aes(x = sex, fill=status)) + geom_bar(position="dodge")



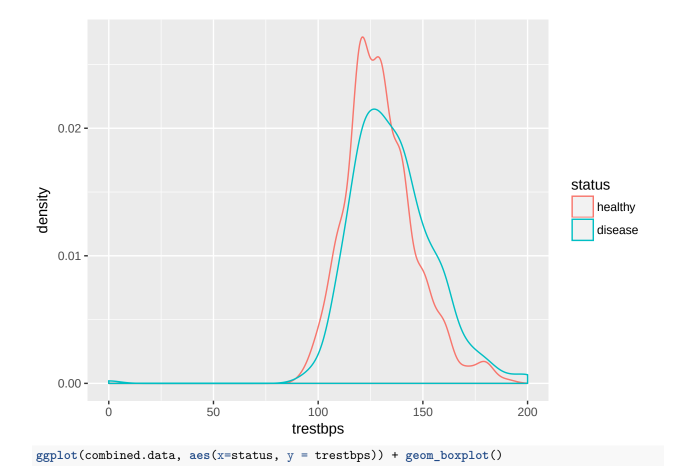


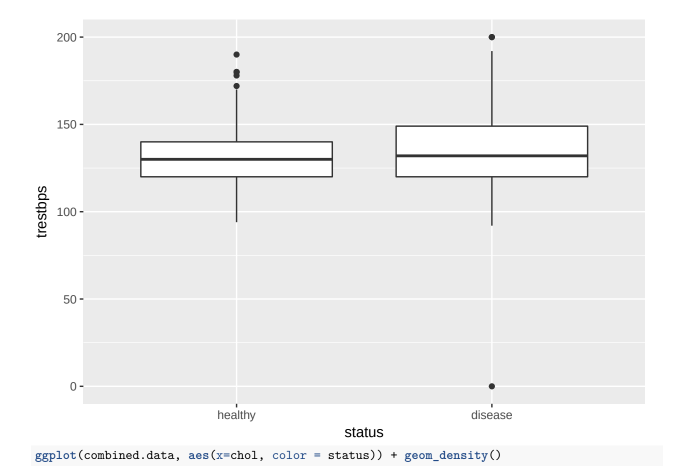


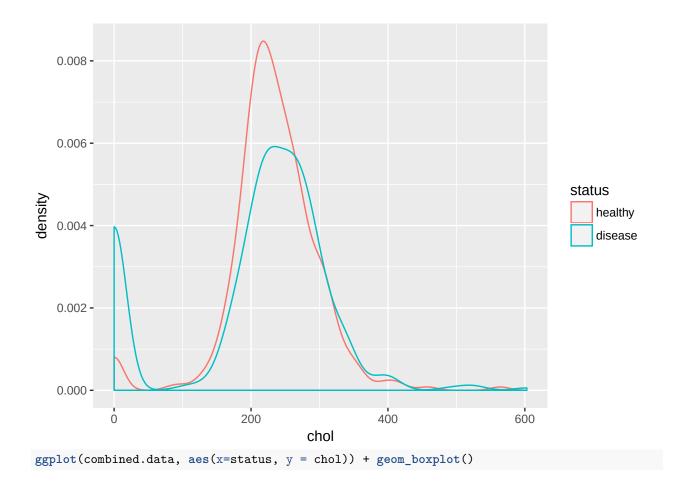
ggplot(combined.data, aes(x = cp, fill=status)) + geom_bar(position="dodge")

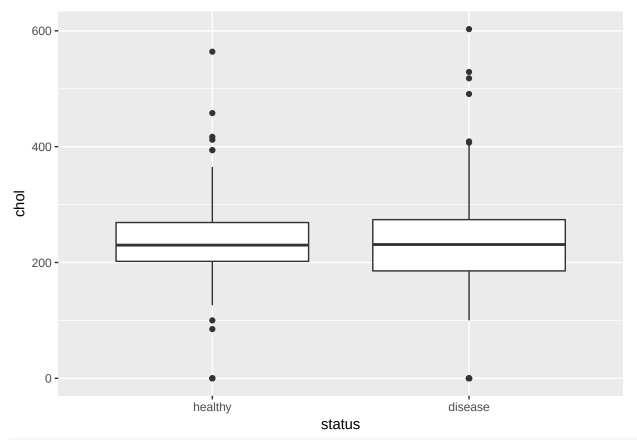


ggplot(combined.data, aes(x=trestbps, color = status)) + geom_density()

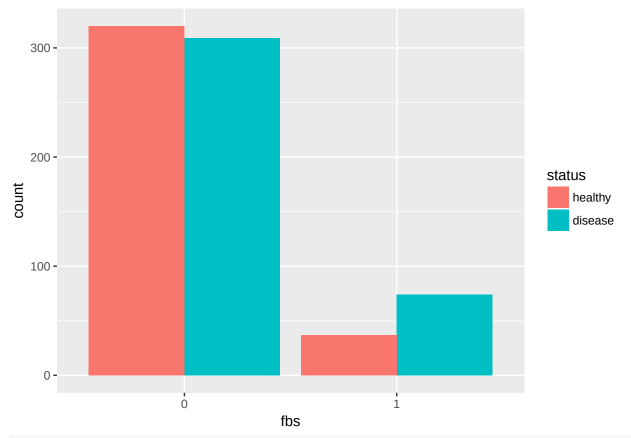




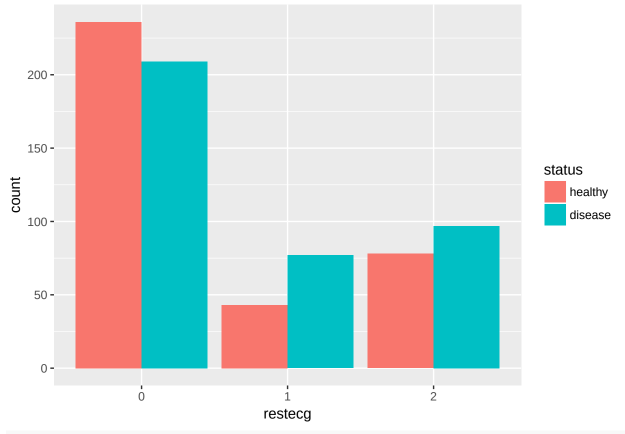




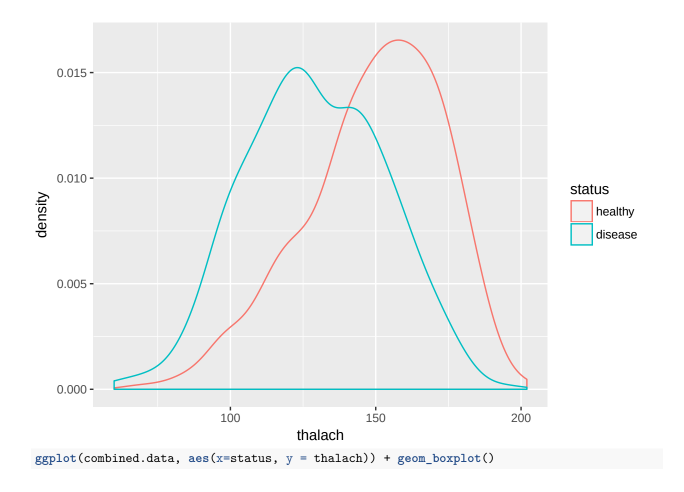
ggplot(combined.data, aes(x = fbs, fill=status)) + geom_bar(position="dodge")

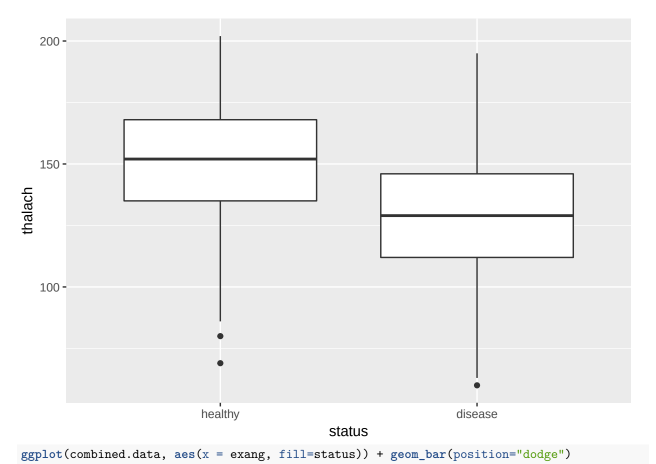


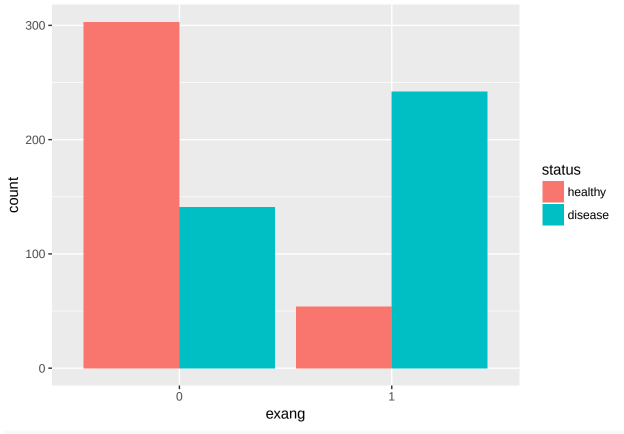
ggplot(combined.data, aes(x = restecg, fill=status)) + geom_bar(position="dodge")



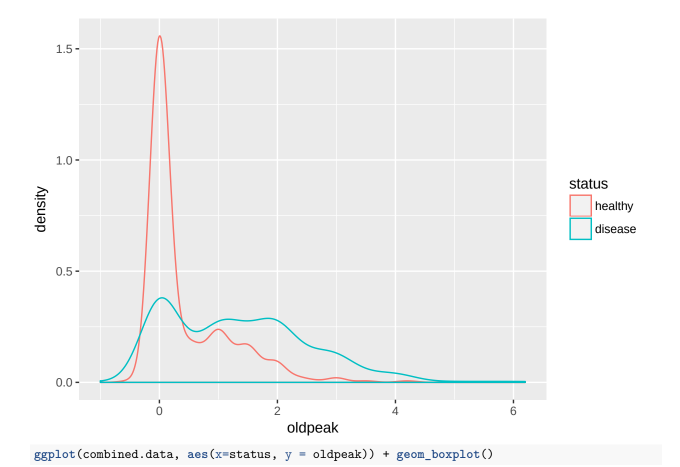
ggplot(combined.data, aes(x=thalach, color = status)) + geom_density()

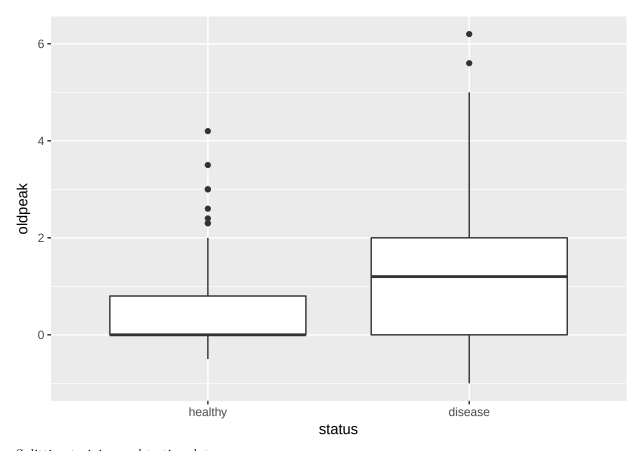






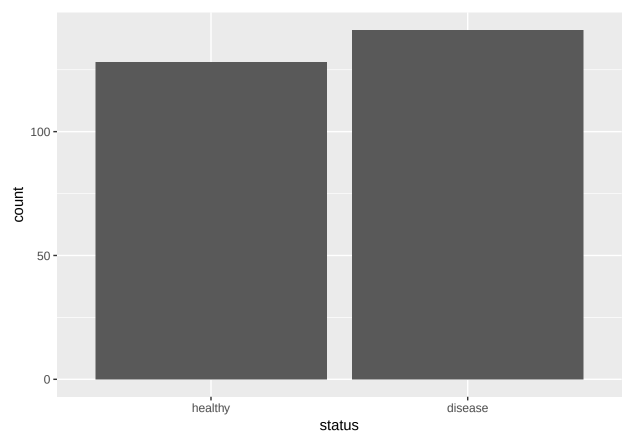
ggplot(combined.data, aes(x=oldpeak, color = status)) + geom_density()





Splitting training and testing data

```
set.seed(100)
sample = sample.split(combined.data, SplitRatio = 0.70)
train = subset(combined.data, sample == T)
test = subset(combined.data, sample == F)
summary(test)
##
                       sex
                                          trestbps
                                                             chol
        age
                                ср
##
   Min. :29.0
                  female: 67
                               1: 14
                                       Min. : 0.0
                                                       Min. : 0.0
   1st Qu.:46.0
                  male :202
                               2: 53
                                        1st Qu.:120.0
                                                       1st Qu.:203.0
##
  Median:54.0
                               3: 59
##
                                       Median :130.0
                                                       Median :234.0
##
  Mean :53.2
                               4:143
                                       Mean
                                             :132.3
                                                       Mean :220.9
   3rd Qu.:60.0
                                        3rd Qu.:140.0
                                                        3rd Qu.:268.0
##
##
  Max.
          :77.0
                                       Max.
                                               :200.0
                                                        Max.
                                                               :564.0
  fbs
##
           restecg
                      thalach
                                               oldpeak
                                                                 status
                                    exang
##
  0:228
           0:166
                   Min. : 69.0
                                   0:164
                                           Min.
                                                   :0.0000
                                                            healthy:128
                                            1st Qu.:0.0000
                   1st Qu.:120.0
##
   1: 41
           1: 44
                                   1:105
                                                            disease:141
##
           2: 59
                   Median :140.0
                                           Median :0.4000
##
                   Mean
                         :139.4
                                           Mean
                                                   :0.8643
##
                   3rd Qu.:160.0
                                            3rd Qu.:1.5000
##
                          :202.0
                                                   :5.0000
                   Max.
                                           Max.
ggplot(test, aes(x = status)) + geom_bar()
```



Baseline approach: Logistic regression (both with full model training, and with the selected five predictors)

```
logit.model = train(status ~ ., data=train, method = 'glm', family = 'binomial')
summary(logit.model)
```

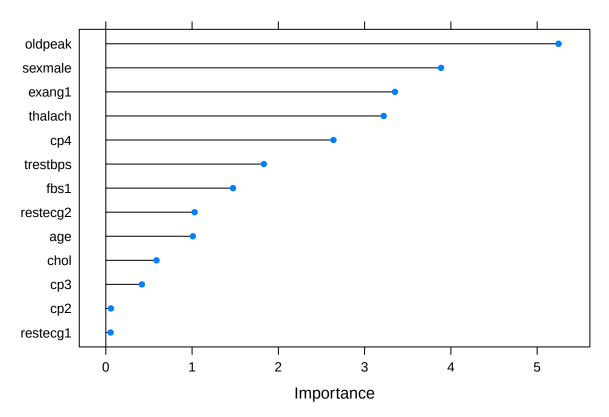
```
##
## Call:
## NULL
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
## -2.8479 -0.6053
                      0.1551
                               0.5699
                                        2.6069
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.9049653 1.7734166 -1.638 0.101409
               0.0162828 0.0161423
                                       1.009 0.313119
## age
## sexmale
                1.3139698
                          0.3380953
                                       3.886 0.000102 ***
               -0.0368386
                           0.6284873
                                      -0.059 0.953259
## cp2
## cp3
                0.2472175
                           0.5918941
                                       0.418 0.676187
## cp4
                1.5137979
                           0.5736822
                                       2.639 0.008321 **
## trestbps
                0.0138848
                           0.0075800
                                       1.832 0.066986
               -0.0009002
                           0.0015325
## chol
                                      -0.587 0.556919
## fbs1
                           0.3725546
                                       1.474 0.140468
                0.5491646
## restecg1
                0.0207829
                           0.3743793
                                       0.056 0.955730
## restecg2
                0.3275127
                           0.3183371
                                       1.029 0.303562
## thalach
               -0.0187948
                           0.0058359
                                      -3.221 0.001279 **
## exang1
               0.9493386 0.2832175
                                       3.352 0.000802 ***
```

```
## oldpeak
               ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 652.59 on 470 degrees of freedom
## Residual deviance: 390.26 on 457 degrees of freedom
## AIC: 418.26
##
## Number of Fisher Scoring iterations: 5
logit.result = predict(logit.model, test)
confusionMatrix(test$status, logit.result)
## Confusion Matrix and Statistics
##
            Reference
##
## Prediction healthy disease
##
     healthy
                 109
                          19
     disease
                         106
##
                  35
##
##
                 Accuracy : 0.7993
##
                   95% CI: (0.7463, 0.8454)
      No Information Rate: 0.5353
##
      P-Value [Acc > NIR] : < 2e-16
##
##
##
                    Kappa: 0.5999
##
  Mcnemar's Test P-Value: 0.04123
##
              Sensitivity: 0.7569
##
##
              Specificity: 0.8480
##
           Pos Pred Value: 0.8516
##
           Neg Pred Value: 0.7518
               Prevalence: 0.5353
##
##
           Detection Rate: 0.4052
##
     Detection Prevalence: 0.4758
##
        Balanced Accuracy: 0.8025
##
##
         'Positive' Class : healthy
smalllogit.model = train(status ~ sex +cp + thalach +exang +oldpeak, data=train, method = 'glm', family
smalllogit.result = predict(smalllogit.model, test)
confusionMatrix(test$status, smalllogit.result)
## Confusion Matrix and Statistics
##
##
            Reference
## Prediction healthy disease
##
     healthy
                 108
                          20
##
     disease
                  32
                         109
##
##
                 Accuracy : 0.8067
```

95% CI: (0.7544, 0.8521)

##

```
No Information Rate: 0.5204
##
       P-Value [Acc > NIR] : <2e-16
##
##
##
                     Kappa : 0.6141
##
   Mcnemar's Test P-Value: 0.1272
##
##
               Sensitivity: 0.7714
               Specificity: 0.8450
##
##
            Pos Pred Value : 0.8438
##
            Neg Pred Value: 0.7730
##
                Prevalence: 0.5204
##
            Detection Rate: 0.4015
##
      Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.8082
##
##
          'Positive' Class : healthy
##
variable importance for logistic model
importance = varImp(logit.model, scale=FALSE)
importance
## glm variable importance
##
            Overall
## oldpeak 5.24782
## sexmale 3.88639
## exang1
            3.35198
## thalach 3.22054
            2.63874
## cp4
## trestbps 1.83177
## fbs1
            1.47405
## restecg2 1.02882
## age
            1.00870
## chol
            0.58742
## cp3
            0.41767
## cp2
            0.05861
## restecg1 0.05551
plot(importance)
```



Approach 2: Naive Bayes classification, with both full model, a small model with 5 predictors, and a "best" model that includes all the predictors except trestbps.

```
naiveBayes.model = naiveBayes(status ~., data = train)
naiveBayes.model
```

```
##
## Naive Bayes Classifier for Discrete Predictors
##
## Call:
## naiveBayes.default(x = X, y = Y, laplace = laplace)
## A-priori probabilities:
## Y
##
     healthy
               disease
## 0.4861996 0.5138004
##
##
  Conditional probabilities:
##
            age
## Y
                  [,1]
                           [,2]
     healthy 50.30131 9.527479
##
     disease 55.63223 8.548681
##
##
##
            sex
## Y
                female
     healthy 0.3580786 0.6419214
##
     disease 0.1033058 0.8966942
##
##
##
            ср
                                   2
## Y
                       1
                                              3
```

```
##
     healthy 0.06550218 0.34934498 0.31441048 0.27074236
##
     disease 0.03305785 0.07024793 0.12396694 0.77272727
##
##
            trestbps
## Y
                  [,1]
                           [,2]
##
     healthy 129.7380 16.77721
##
     disease 136.0744 18.21457
##
##
            chol
## Y
                  [,1]
                           [,2]
##
     healthy 230.5939 65.9265
     disease 209.3430 109.4753
##
##
##
            fbs
## Y
                      0
##
     healthy 0.8908297 0.1091703
##
     disease 0.8140496 0.1859504
##
##
            restecg
## Y
##
     healthy 0.6593886 0.1179039 0.2227074
##
     disease 0.5289256 0.2024793 0.2685950
##
##
            thalach
## Y
                  [,1]
                           [,2]
##
     healthy 149.2314 22.97049
##
     disease 128.0372 23.89766
##
##
            exang
## Y
##
     healthy 0.8384279 0.1615721
     disease 0.3636364 0.6363636
##
##
##
            oldpeak
## Y
                   [,1]
     healthy 0.4039301 0.6572352
##
##
     disease 1.3917355 1.2229538
naiveBayes.result = predict(naiveBayes.model, train)
confusionMatrix(train$status, naiveBayes.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
##
      healthy
                   188
                            41
##
      disease
                   47
                           195
##
                  Accuracy : 0.8132
##
##
                    95% CI: (0.775, 0.8474)
##
       No Information Rate: 0.5011
##
       P-Value [Acc > NIR] : <2e-16
##
##
                      Kappa: 0.6263
    Mcnemar's Test P-Value: 0.594
```

```
##
##
               Sensitivity: 0.8000
##
               Specificity: 0.8263
##
            Pos Pred Value: 0.8210
##
            Neg Pred Value: 0.8058
##
                Prevalence: 0.4989
##
            Detection Rate: 0.3992
      Detection Prevalence: 0.4862
##
##
         Balanced Accuracy: 0.8131
##
##
          'Positive' Class : healthy
##
smallnaiveBayes.model = naiveBayes(status ~sex +cp + thalach +exang +oldpeak, data = train)
smallnaiveBayes.result = predict(smallnaiveBayes.model, test)
confusionMatrix(test$status, smallnaiveBayes.result)
## Confusion Matrix and Statistics
##
             Reference
## Prediction healthy disease
                           20
##
     healthy
                  108
##
      disease
                   35
                          106
##
##
                  Accuracy: 0.7955
                    95% CI : (0.7423, 0.8421)
##
##
       No Information Rate: 0.5316
##
       P-Value [Acc > NIR] : < 2e-16
##
##
                     Kappa: 0.5923
  Mcnemar's Test P-Value: 0.05906
##
##
##
               Sensitivity: 0.7552
##
               Specificity: 0.8413
            Pos Pred Value: 0.8438
##
##
            Neg Pred Value: 0.7518
                Prevalence: 0.5316
##
##
            Detection Rate: 0.4015
##
     Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.7983
##
##
          'Positive' Class : healthy
##
bestnaiveBayes.model = naiveBayes(status ~sex +cp + thalach +exang +oldpeak+restecg+fbs+chol+age, data
bestnaiveBayes.result = predict(bestnaiveBayes.model, test)
confusionMatrix(test$status, bestnaiveBayes.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
                  109
##
     healthy
                           19
```

##

##

disease

24

117

```
##
                  Accuracy : 0.8401
##
                    95% CI: (0.7908, 0.8818)
       No Information Rate: 0.5056
##
       P-Value [Acc > NIR] : <2e-16
##
##
##
                     Kappa: 0.6801
   Mcnemar's Test P-Value: 0.5419
##
##
##
               Sensitivity: 0.8195
##
               Specificity: 0.8603
##
            Pos Pred Value: 0.8516
##
            Neg Pred Value: 0.8298
##
                Prevalence: 0.4944
##
            Detection Rate: 0.4052
##
      Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.8399
##
##
          'Positive' Class : healthy
##
```

Approach 3: Support vector machines, using a 3-repeat, 10-fold cross validation to fine tune the hyperparameters C, using Gaussian radial kernel. Includes both the results of the full model and a small model with the selected 5 predictors.

```
fitControl = trainControl(method = "repeatedcv", number = 10, repeats = 3)
set.seed(100)
svm.model = train(status ~., data = train, method = "svmRadial", trControl = fitControl)
svm.model
## Support Vector Machines with Radial Basis Function Kernel
##
## 471 samples
  10 predictor
##
    2 classes: 'healthy', 'disease'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 424, 424, 424, 424, 424, 424, ...
## Resampling results across tuning parameters:
##
##
           Accuracy
                      Kappa
##
     0.25 0.7890052 0.5774449
##
     0.50 0.7911630 0.5825566
##
     1.00 0.7926425 0.5856137
##
## Tuning parameter 'sigma' was held constant at a value of 0.05299966
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were sigma = 0.05299966 and C = 1.
svm.result = predict(svm.model, newdata = test)
confusionMatrix(test$status, svm.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
```

```
##
      healthy
                  111
                           17
                          108
##
      disease
                   33
##
##
                  Accuracy : 0.8141
##
                    95% CI: (0.7624, 0.8588)
##
       No Information Rate: 0.5353
##
       P-Value [Acc > NIR] : < 2e-16
##
##
                     Kappa: 0.6295
##
   Mcnemar's Test P-Value: 0.03389
##
##
               Sensitivity: 0.7708
##
               Specificity: 0.8640
##
            Pos Pred Value: 0.8672
##
            Neg Pred Value: 0.7660
##
                Prevalence: 0.5353
##
            Detection Rate: 0.4126
##
      Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.8174
##
##
          'Positive' Class : healthy
##
set.seed(100)
smallsvm.model = train(status ~sex +cp + thalach +exang +oldpeak, data = train,
                 method = "svmRadial",
                 trControl = fitControl)
smallsvm.result = predict(smallsvm.model, newdata = test)
confusionMatrix(test$status, smallsvm.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
##
      healthy
                  112
                           16
##
      disease
                   41
                          100
##
##
                  Accuracy: 0.7881
                    95% CI: (0.7344, 0.8354)
##
##
       No Information Rate: 0.5688
##
       P-Value [Acc > NIR] : 3.319e-14
##
##
                     Kappa: 0.579
##
   Mcnemar's Test P-Value: 0.001478
##
##
               Sensitivity: 0.7320
##
               Specificity: 0.8621
##
            Pos Pred Value: 0.8750
            Neg Pred Value: 0.7092
##
##
                Prevalence: 0.5688
##
            Detection Rate: 0.4164
##
      Detection Prevalence: 0.4758
         Balanced Accuracy: 0.7970
##
##
##
          'Positive' Class : healthy
```

##

Variable importance for the full sym model.

```
importance = varImp(svm.model, scale=FALSE)
importance
## ROC curve variable importance
##
##
            Importance
                 0.7570
## ср
## oldpeak
                 0.7479
## thalach
                 0.7430
                 0.7374
## exang
## age
                 0.6689
## sex
                 0.6274
## trestbps
                 0.6037
## restecg
                 0.5585
## fbs
                 0.5384
## chol
                 0.5180
plot(importance)
     ср
oldpeak
thalach
 exang
    age
    sex
trestbps
restecg
    fbs
   chol
```

Approach 4: Random forests, with 10 fold, 3 repeat cross validation to tune the parameter mtry. With results from both the full model and the small model using the 5 selected predictors

Importance

0.65

0.70

0.75

0.60

```
fitControl = trainControl(method = "repeatedcv", number = 10, repeats = 3)
set.seed(100)
rf.model = train(status~., data=train, method="rf", trControl = fitControl)
rf.model
```

Random Forest

0.55

```
##
## 471 samples
##
    10 predictor
     2 classes: 'healthy', 'disease'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 424, 424, 424, 424, 424, 424, ...
## Resampling results across tuning parameters:
##
##
     mtry
           Accuracy
                      Kappa
      2
##
           0.7855053
                      0.5706418
      7
                      0.5366725
##
           0.7685130
           0.7550667
                      0.5098978
##
     13
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
rf.result = predict(rf.model, test)
confusionMatrix(test$status, rf.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
##
      healthy
                  109
                            19
##
      disease
                   30
                           111
##
##
                  Accuracy: 0.8178
##
                    95% CI: (0.7664, 0.8621)
##
       No Information Rate: 0.5167
       P-Value [Acc > NIR] : <2e-16
##
##
##
                     Kappa: 0.6363
##
    Mcnemar's Test P-Value: 0.1531
##
##
               Sensitivity: 0.7842
               Specificity: 0.8538
##
##
            Pos Pred Value: 0.8516
##
            Neg Pred Value: 0.7872
##
                Prevalence: 0.5167
##
            Detection Rate: 0.4052
      Detection Prevalence : 0.4758
##
##
         Balanced Accuracy: 0.8190
##
##
          'Positive' Class : healthy
##
predictors(rf.model)
                   "sexmale"
    [1] "age"
                               "cp2"
                                          "cp3"
                                                      "cp4"
                                                                 "trestbps"
                               "restecg1" "restecg2" "thalach"
##
    [7] "chol"
                   "fbs1"
                                                                 "exang1"
## [13] "oldpeak"
```

Small random forests model using only the 5 selected predictors.

```
set.seed(100)
fitControl = trainControl(method = "repeatedcv", number = 10, repeats = 3)
smallrf.model = train(status~sex +cp + thalach +exang +oldpeak, data=train, method="rf")
## Random Forest
##
## 471 samples
##
     5 predictor
     2 classes: 'healthy', 'disease'
##
## No pre-processing
## Resampling: Bootstrapped (25 reps)
## Summary of sample sizes: 471, 471, 471, 471, 471, 471, ...
## Resampling results across tuning parameters:
##
##
     mtry Accuracy
                      Kappa
##
           0.7731039 0.5455893
     2
##
           0.7484221
                      0.4966682
##
           0.7363129 0.4727022
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
smallrf.result = predict(smallrf.model, test)
confusionMatrix(test$status, smallrf.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
##
      healthy
                  107
                           21
##
      disease
                   36
                          105
##
##
                  Accuracy : 0.7881
##
                    95% CI: (0.7344, 0.8354)
       No Information Rate: 0.5316
##
       P-Value [Acc > NIR] : < 2e-16
##
##
##
                     Kappa: 0.5775
##
   Mcnemar's Test P-Value: 0.06369
##
               Sensitivity: 0.7483
##
##
               Specificity: 0.8333
            Pos Pred Value: 0.8359
##
##
            Neg Pred Value: 0.7447
##
                Prevalence: 0.5316
            Detection Rate: 0.3978
##
##
      Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.7908
##
##
          'Positive' Class : healthy
```

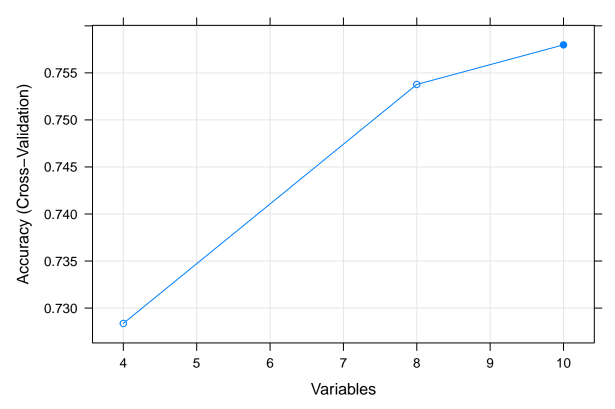
Variable importance for the full random forest model

```
importance = varImp(rf.model, scale=FALSE)
importance
## rf variable importance
##
##
            Overall
## thalach
             29.115
## oldpeak
             27.043
             22.971
## cp4
             20.156
## age
## exang1
             19.875
## chol
             19.073
## trestbps 16.695
## sexmale
              9.497
              9.348
## cp2
## cp3
              5.525
## fbs1
              4.020
## restecg1
              3.821
## restecg2
              3.149
plot(importance)
 thalach
oldpeak
    ср4
    age
 exang1
    chol
trestbps
sexmale
    cp2
    ср3
   fbs1
restecg1
restecg2
                 5
                              10
                                           15
                                                         20
                                                                      25
                                                                                   30
                                         Importance
```

Variable selection via recursive variable elimination using random forests

```
set.seed(100)
control = rfeControl(functions=rfFuncs, method="cv", number=10, repeats = 3, returnResamp = "all")
results = rfe(train[, 1:10], train[, 11], rfeControl=control)
results
##
## Recursive feature selection
```

```
##
## Outer resampling method: Cross-Validated (10 fold)
## Resampling performance over subset size:
##
## Variables Accuracy Kappa AccuracySD KappaSD Selected
           4 0.7284 0.4564 0.06860 0.1375
           8 0.7538 0.5078
                                0.05571 0.1113
##
##
          10 0.7580 0.5162
                              0.06024 0.1197
##
## The top 5 variables (out of 10):
      cp, oldpeak, exang, sex, thalach
print(results)
##
## Recursive feature selection
## Outer resampling method: Cross-Validated (10 fold)
## Resampling performance over subset size:
##
## Variables Accuracy Kappa AccuracySD KappaSD Selected
##
           4 0.7284 0.4564 0.06860 0.1375
##
           8
              0.7538 0.5078
                                0.05571 0.1113
              0.7580 0.5162
                                0.06024 0.1197
##
          10
##
## The top 5 variables (out of 10):
      cp, oldpeak, exang, sex, thalach
predictors(results)
## [1] "cp"
                  "oldpeak"
                             "exang"
                                        "sex"
                                                   "thalach" "age"
## [7] "fbs"
                  "chol"
                             "restecg" "trestbps"
plot(results, type = c("g", "o"))
```



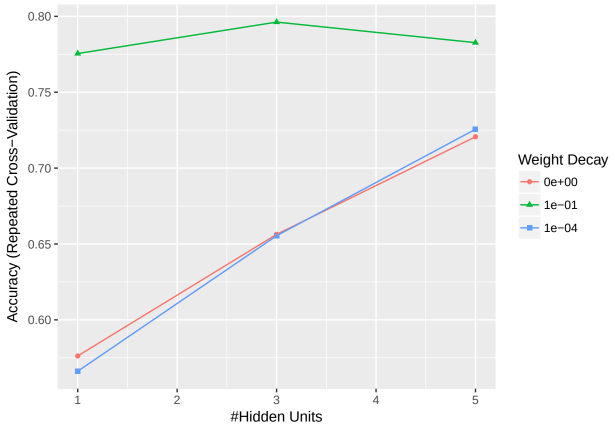
Approach 5: Single layer neural network, using 10 fold, 3 repeat cross validation to tune the parameters size and decay.

```
fitControl = trainControl(method = "repeatedcv", number = 10, repeats = 3)
set.seed(100)
nnet.model = train(status ~ ., data = train, method = "nnet", trControl = fitControl, verbose = FALSE,
nnet.model
## Neural Network
##
## 471 samples
   10 predictor
##
     2 classes: 'healthy', 'disease'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 424, 424, 424, 424, 424, 424, ...
## Resampling results across tuning parameters:
##
##
           decay Accuracy
                             Kappa
##
           0e+00
                  0.5762129
                             0.1388614
     1
                  0.5661360 0.1161913
##
     1
           1e-04
##
           1e-01
                  0.7755011 0.5522494
     1
##
     3
           0e+00
                  0.6563483
                             0.3152133
##
     3
           1e-04
                  0.6554091
                             0.3066531
##
     3
           1e-01
                  0.7962181
                             0.5927021
##
     5
           0e+00
                  0.7206413
                             0.4429703
##
     5
           1e-04
                  0.7256090
                             0.4526095
                  0.7826678 0.5655450
##
     5
           1e-01
##
```

```
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were size = 3 and decay = 0.1.
nnet.result = predict(nnet.model, test)
confusionMatrix(test$status, nnet.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
                  113
##
      healthy
                           15
      disease
                          104
##
                   37
##
##
                  Accuracy: 0.8067
##
                    95% CI : (0.7544, 0.8521)
       No Information Rate: 0.5576
##
       P-Value [Acc > NIR] : < 2.2e-16
##
##
##
                     Kappa: 0.6155
##
   Mcnemar's Test P-Value: 0.003589
##
##
               Sensitivity: 0.7533
##
               Specificity: 0.8739
##
            Pos Pred Value: 0.8828
##
            Neg Pred Value: 0.7376
##
                Prevalence: 0.5576
##
            Detection Rate: 0.4201
      Detection Prevalence: 0.4758
##
##
         Balanced Accuracy: 0.8136
##
##
          'Positive' Class : healthy
##
set.seed(100)
smallnnet.model = train(status ~ sex +cp + thalach +exang +oldpeak, data = train, method = "nnet",
smallnnet.model
## Neural Network
##
## 471 samples
##
     5 predictor
     2 classes: 'healthy', 'disease'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold, repeated 3 times)
## Summary of sample sizes: 424, 424, 424, 424, 424, 424, ...
## Resampling results across tuning parameters:
##
##
     size
           decay Accuracy
                             Kappa
##
           0e+00
                  0.5329357 0.04123052
           1e-04 0.5230066 0.01914251
##
##
           1e-01 0.7770981 0.55263622
     1
##
           0e+00 0.5857308 0.15125408
     3
##
     3
           1e-04
                  0.6149688 0.21190317
##
     3
           1e-01 0.7875591 0.57553217
##
     5
           0e+00 0.5984794 0.17879022
```

```
##
           1e-04 0.6477696 0.28080780
##
    5
           1e-01 0.7854925 0.57149119
##
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were size = 3 and decay = 0.1.
smallnnet.result = predict(smallnnet.model, test)
confusionMatrix(test$status, smallnnet.result)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction healthy disease
      healthy
                  111
                           17
##
      disease
                   46
                           95
##
##
                  Accuracy : 0.7658
                    95% CI : (0.7106, 0.8151)
##
##
       No Information Rate: 0.5836
##
       P-Value [Acc > NIR] : 2.732e-10
##
##
                     Kappa: 0.5354
   Mcnemar's Test P-Value: 0.0004192
##
##
##
               Sensitivity: 0.7070
##
               Specificity: 0.8482
##
            Pos Pred Value: 0.8672
            Neg Pred Value: 0.6738
##
##
                Prevalence: 0.5836
##
            Detection Rate: 0.4126
##
      Detection Prevalence: 0.4758
##
         Balanced Accuracy: 0.7776
##
##
          'Positive' Class : healthy
variable importance based on neural network model
ggplot(nnet.model)
```

Warning: Ignoring unknown aesthetics: shape



```
importance = varImp(nnet.model, scale=FALSE)
importance
```

```
## nnet variable importance
##
            Overall
##
## cp2
            15.4286
## exang1
            14.2669
            12.4885
## cp4
## cp3
            12.3490
## oldpeak 10.5689
## restecg1 10.1935
## sexmale
             9.9430
## fbs1
             7.8338
## restecg2 2.7411
             2.0853
## age
## trestbps 1.0302
## chol
             0.8389
## thalach
             0.2322
plot(importance)
```

