# 8106final code

# Ze Li

# load libraries
library(dplyr)

train <- as.data.frame(training\_data)
test <- as.data.frame(test\_data)</pre>

head(train)

```
library(tidyverse)
library(corrplot)
library(gridExtra)
library(ggplot2)
library(patchwork)
library(MASS)
library(mgcv)
library(earth)
library(Formula)
library(plotmo)
library(plotrix)
library(TeachingDemos)
library(caret)
library(glmnet)
library(tidymodels)
library(mlbench)
library(pROC)
library(pdp)
library(vip)
library(AppliedPredictiveModeling)
library(rsample)
library(klaR)
library(rpart)
library(rpart.plot)
library(party)
library(partykit)
library(randomForest)
library(ranger)
library(gbm)
library(e1071)
library(kernlab)
library(ggrepel)
# load data
load("/Users/zeze/Library/Mobile Documents/com~apple~CloudDocs/2024/24S BIST P8106 DS II/8106final/p810
load("/Users/zeze/Library/Mobile Documents/com~apple~CloudDocs/2024/24S BIST P8106 DS II/8106final/p810
```

```
id age gender race smoking height weight bmi diabetes hypertension SBP LDL
## 1 1 59
                0
                      1
                              1 170.3
                                        74.7 25.8
                                                         0
                                                                      0 120
## 2 2
                              1 170.8
                                        75.7 26.0
        54
                1
                     1
                                                                      1 133 87
## 3 3 55
                     3
                                172.7
                                        89.5 30.0
                                                         0
                                                                      0 123 139
                1
                              1
## 4 4
        59
                0
                     1
                                171.7
                                        74.6 25.3
                                                         0
                                                                      0 121 126
## 6 6
        64
                1
                     1
                              0 168.8
                                       87.9 30.8
                                                         0
                                                                      1 132 99
## 9 9 67
                0
                     1
                              0 168.5
                                       76.5 27.0
                                                         0
                                                                      1 138 97
     vaccine depression severity
## 1
          1
                     5
                               0
## 2
          0
                     2
                               1
## 3
          1
                     5
                               0
## 4
                     4
                               0
          1
                     9
                               0
## 6
          1
                     8
## 9
          0
                               1
```

#### head(test)

```
##
      id age gender race smoking height weight bmi diabetes hypertension SBP LDL
## 5
                               0 160.2
                                          75.8 29.5
      5 62
                 1
                       1
                                                           1
                                                                        0 122 107
                               0 172.4
## 7
      7 64
                                          80.9 27.2
                                                           0
                                                                        0 122 99
                  0
                       1
## 8
         62
                       4
                               1 175.6
                                          77.4 25.1
                                                           0
                                                                        0 119 123
      8
                 1
                               0 174.7
## 16 16 62
                 1
                       1
                                          92.4 30.3
                                                           0
                                                                        1 146 108
## 20 20 56
                               0 172.3
                 0
                       1
                                         74.2 25.0
                                                           0
                                                                        0 130 146
## 23 23 66
                  1
                       1
                               1 173.2
                                          68.3 22.8
                                                           0
                                                                       1 133 106
##
      vaccine depression severity
## 5
            1
                       9
## 7
                       7
           0
                                0
## 8
           0
                       6
                                0
## 16
           0
                       4
                                1
## 20
                       7
           1
                                0
## 23
                                1
```

```
train.raw =
 train[, -1] %>%
  mutate(
    gender = as.factor(gender),
   race = as.factor(race),
   smoking = as.factor(smoking),
   hypertension = as.factor(hypertension),
   diabetes = as.factor(diabetes),
   vaccine = as.factor(vaccine),
   severity = as.factor(severity)
test.raw =
 test[, -1] %>%
 mutate(
   gender = as.factor(gender),
   race = as.factor(race),
   smoking = as.factor(smoking),
   hypertension = as.factor(hypertension),
   diabetes = as.factor(diabetes),
   vaccine = as.factor(vaccine),
```

```
severity = as.factor(severity)
)

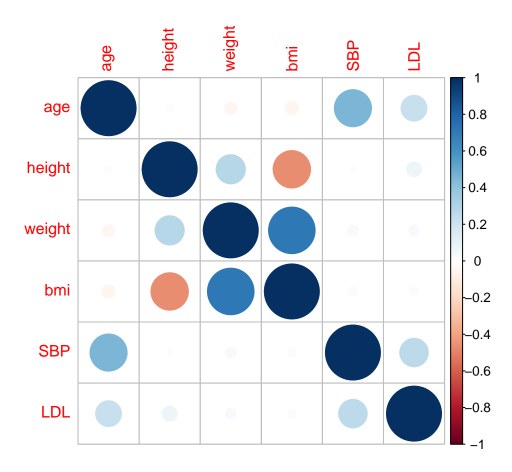
x_train <- model.matrix(severity ~ . , train.raw)[, -1]
y_train <- train.raw$severity
x_test <- model.matrix(severity ~ . , test.raw)[, -1]
y_test <- test.raw$severity
head(train.raw)</pre>
```

```
age gender race smoking height weight bmi diabetes hypertension SBP LDL
## 1 59 0 1
                   1 170.3 74.7 25.8
                                            0
                                                          0 120 95
## 2 54 1 1
## 3 55 1 3
                                             1
                      1 170.8 75.7 26.0
                                                          1 133 87
                     1 172.7 89.5 30.0 0
0 171.7 74.6 25.3 0
0 168.8 87.9 30.8 0
0 168.5 76.5 27.0 0
                     1 172.7 89.5 30.0
                                                         0 123 139
## 4 59
          0 1
                                                         0 121 126
## 6 64
           1 1
                                                         1 132 99
## 9 67 0
              1
                                                         1 138 97
## vaccine depression severity
## 1
      1 5
                          0
                2
5
## 2
        0
                          1
## 3
       1
                          0
## 4
        1
                 4
                          0
## 6
## 9
       1
                 9
                          0
                  8
        0
```

# **Exploratary Data Analysis**

## Correlation plot

```
corr_dat = train.raw %>%
  dplyr::select('age', 'height', 'weight', 'bmi', 'SBP', 'LDL')
corrplot(cor(corr_dat), method = "circle", type = "full")
```

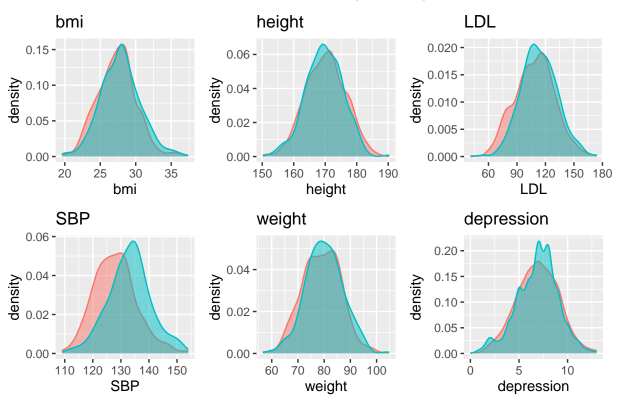


## Continuous Variables

```
p_bmi <- ggplot(train.raw, aes(x = bmi, fill = as.factor(severity),</pre>
                                color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "bmi") +
  theme(legend.position = "none")
p_height <- ggplot(train.raw, aes(x = height, fill = as.factor(severity),</pre>
                                   color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "height") +
  theme(legend.position = "none")
p_ldl <- ggplot(train.raw, aes(x = LDL, fill = as.factor(severity),</pre>
                                color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "LDL") +
  theme(legend.position = "none")
p_sbp <- ggplot(train.raw, aes(x = SBP, fill = as.factor(severity),</pre>
                                color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "SBP") +
```

```
theme(legend.position = "none")
p_weight <- ggplot(train.raw, aes(x = weight, fill = as.factor(severity),</pre>
                                   color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "weight") +
  theme(legend.position = "none")
p_depression <- ggplot(train.raw, aes(x = depression, fill = as.factor(severity),</pre>
                                       color = as.factor(severity))) +
  geom_density(alpha = 0.5) +
  labs(title = "depression") +
  theme(legend.position = "none")
plot_grid <- p_bmi + p_height + p_ldl + p_sbp + p_weight + p_depression +</pre>
  plot_layout(ncol = 3, byrow = TRUE) +
  plot_annotation(title="Continuous variables by Severity", # Adding the title here
                  theme = theme(plot.title = element_text(hjust = 0.5))) # Center the title)
plot_grid
```

# Continuous variables by Severity

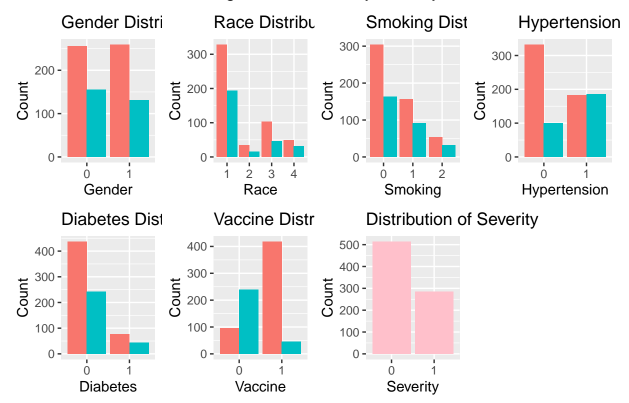


### Categorical variables

```
# Bar Plot for Gender
```

```
p11 <- ggplot(train.raw, aes(x = as.factor(gender), fill = as.factor(severity))) +
  geom_bar(position = "dodge") +
  labs(title = "Gender Distribution", x = "Gender", y = "Count") +
  theme(legend.position = "none")
# Bar Plot for Race
p12 <- ggplot(train.raw, aes(x = as.factor(race), fill = as.factor(severity))) +
  geom_bar(position = "dodge") +
  labs(title = "Race Distribution", x = "Race", y = "Count") +
  theme(legend.position = "none")
# Bar Plot for Smoking Status
p13 <- ggplot(train.raw, aes(x = as.factor(smoking), fill = as.factor(severity))) +
  geom_bar(position = "dodge") +
  labs(title = "Smoking Distribution", x = "Smoking", y = "Count") +
  theme(legend.position = "none")
# Bar Plot for Hypertension
p14 <- ggplot(train.raw, aes(x = as.factor(hypertension), fill = as.factor(severity))) +
  geom_bar(position = "dodge") +
  labs(title = "Hypertension Distribution", x = "Hypertension", y = "Count") +
  theme(legend.position = "none")
# Bar Plot for Diabetes
p15 <- ggplot(train.raw, aes(x = as.factor(diabetes), fill = as.factor(severity))) +
  geom bar(position = "dodge") +
  labs(title = "Diabetes Distribution", x = "Diabetes", y = "Count") +
  theme(legend.position = "none")
# Bar plot for Vaccine
p16 <- ggplot(train.raw, aes(x = as.factor(vaccine), fill = as.factor(severity))) +</pre>
  geom_bar(position = "dodge") +
  labs(title = "Vaccine Distribution", x = "Vaccine", y = "Count") +
  theme(legend.position = "none")
# Bar plot for Severity
p17 <- ggplot(train.raw, aes(x = severity)) +
  geom bar(fill = "#FFCOCB") +
  labs(title = "Distribution of Severity", x = "Severity", y = "Count")
# Combine the plots into a 2x4 grid
plot_grid2 <- p11 + p12 + p13 + p14 + p15 + p16 + p17 +
  plot_layout(ncol = 4, byrow = TRUE) +
  plot_annotation(title="Categorical variables by Severity", # Adding the title here
                  theme = theme(plot.title = element_text(hjust = 0.5))) # Center the title)
# Display the combined plot
plot_grid2
```

# Categorical variables by Severity



# Model fitting

logistic regression

```
glm
##
## Call:
## NULL
##
```

```
## (Intercept) -36.142676 30.259726 -1.194 0.23232
                         0.028105 2.305 0.02114 *
               0.064795
## age
## gender1
               -0.409132
                         0.211384 -1.935 0.05293
## race2
               -0.202620 0.441920 -0.458 0.64659
## race3
               0.017372 0.288969 0.060 0.95206
## race4
               -0.174620 0.352767 -0.495 0.62060
               0.024966 0.234778 0.106 0.91531
## smoking1
## smoking2
               0.492400 0.352148 1.398 0.16203
## height
               ## weight
               ## bmi
                ## diabetes1
## hypertension1 0.380927
                          0.340032 1.120 0.26260
## SBP
                0.070811
                          0.023346
                                  3.033 0.00242 **
## LDL
                          0.005828
                                  1.720 0.08548 .
                0.010022
## vaccine1
               -3.617987
                          0.241871 -14.958 < 2e-16 ***
## depression
               -0.037969 0.050059 -0.758 0.44816
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 1043.15 on 799 degrees of freedom
## Residual deviance: 593.41 on 783 degrees of freedom
## AIC: 627.41
## Number of Fisher Scoring iterations: 5
## test error
glm.pred <- predict(model.glm, newdata = x_test, type = "prob")[,2]</pre>
roc.glm <- roc(y_test, glm.pred)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases
1-roc.glm$auc[1]
## [1] 0.1039316
glmnGrid <- expand.grid(.alpha = seq(0, 1, length = 21),</pre>
                     .lambda = exp(seq(-5, 5, length = 50)))
set.seed(83)
model.glmn <- train(x = x_train,</pre>
                 y = y_train,
                 method = "glmnet",
                 tuneGrid = glmnGrid,
```

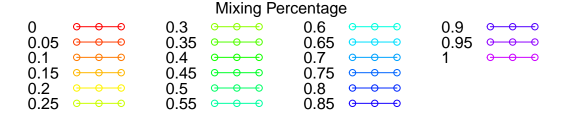
Estimate Std. Error z value Pr(>|z|)

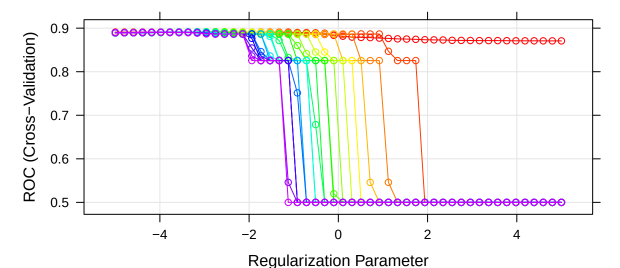
## Coefficients:

##

## Penalized logistic regression

```
## alpha lambda
## 263 0.25 0.07800203
```





```
# test error
glmn.pred <- predict(model.glmn, newdata = x_test, type = "prob")[,2]
roc.glmn <- roc(y_test, glmn.pred)</pre>
```

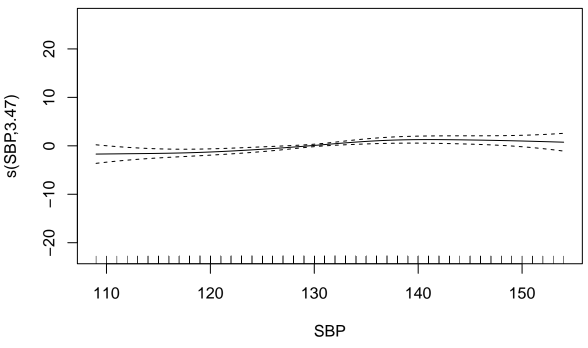
```
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases</pre>
```

```
1-roc.glmn$auc[1]
```

## [1] 0.1039316

#### GAM

```
##
## Family: binomial
## Link function: logit
##
## Formula:
## .outcome ~ gender1 + race2 + race3 + race4 + smoking1 + smoking2 +
##
       diabetes1 + hypertension1 + vaccine1 + s(depression) + s(age) +
       s(SBP) + s(LDL) + s(bmi) + s(height) + s(weight)
##
##
## Estimated degrees of freedom:
## 1.00 1.06 3.47 1.74 1.00 7.69 1.00
## total = 26.97
##
## UBRE score: -0.2345662
plot(model.gam$finalModel, select = 3)
```



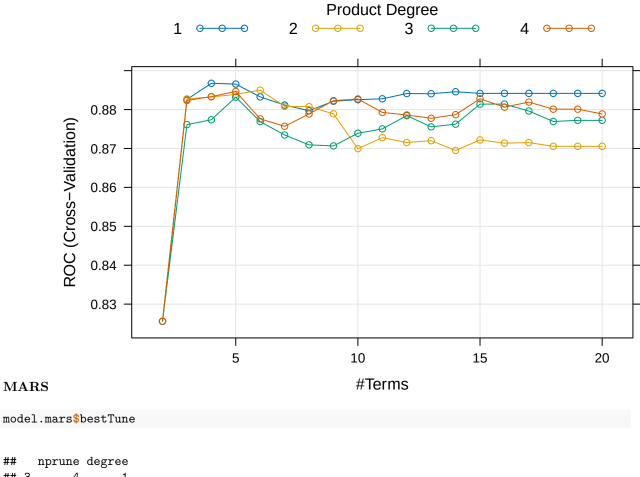
```
# test error
gam.pred <- predict(model.gam, newdata = x_test, type = "prob")[,2]
roc.gam <- roc(y_test, gam.pred)

## Setting levels: control = 0, case = 1

## Setting direction: controls < cases

1-roc.gam$auc[1]</pre>
```

## [1] 0.102792

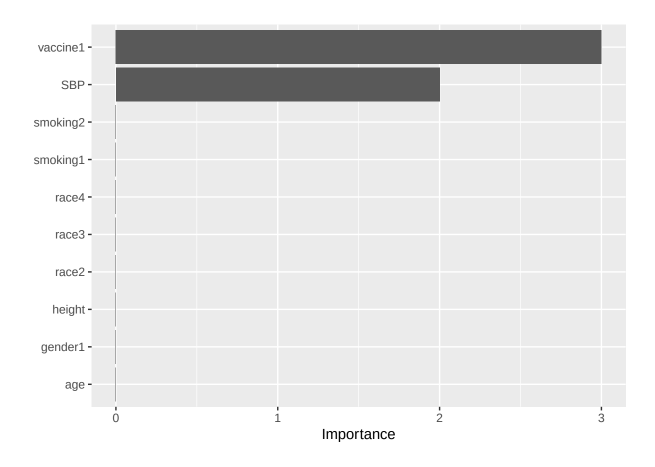


```
## 3
          4
```

```
coef(model.mars$finalModel)
```

```
## (Intercept)
                   vaccine1
                              h(SBP-139)
                                         h(139-SBP)
   2.193120089 -3.323844997 -0.002886632 -0.129783879
```

```
#pdp::partial(model.mars, pred.var = c("age"), grid.resolution = 200) %>% autoplot()
vip(model.mars$finalModel, type = "nsubsets")
```



```
# test error
glm.pred <- predict(model.glm, newdata = x_test, type = "prob")[,2]
glmn.pred <- predict(model.glmn, newdata = x_test, type = "prob")[,2]
gam.pred <- predict(model.gam, newdata = x_test, type = "prob")[,2]
mars.pred <- predict(model.mars, newdata = x_test, type = "prob")[,2]
roc.glm <- roc(y_test, glm.pred)</pre>
```

## logistic regression test performance

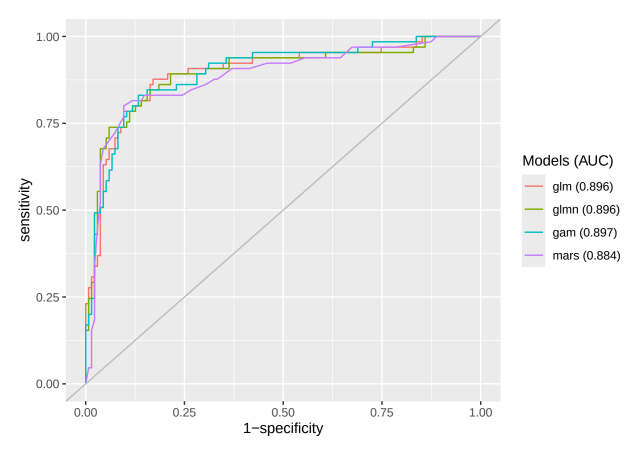
```
## Setting levels: control = 0, case = 1
## Setting direction: controls < cases

roc.glmn <- roc(y_test, glmn.pred)

## Setting levels: control = 0, case = 1
## Setting direction: controls < cases

roc.gam <- roc(y_test, gam.pred)

## Setting levels: control = 0, case = 1
## Setting direction: controls < cases</pre>
```



## LDA

## QDA

```
## X0 X1

## 5 0.98826264 0.01173736

## 7 0.11045382 0.88954618

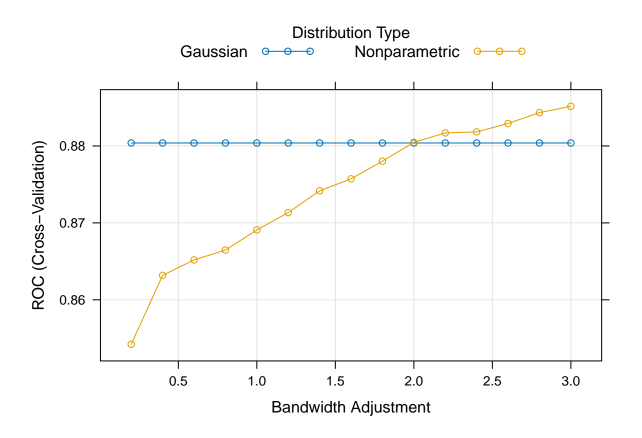
## 8 0.43666109 0.56333891

## 16 0.02962259 0.97037741

## 20 0.97869427 0.02130573

## 23 0.06623623 0.93376377
```

## naive bayers NB



```
# test performance
roc.lda <- roc(y_test, lda.pred2[,2])

lda qda test perforance

## Setting levels: control = 0, case = 1

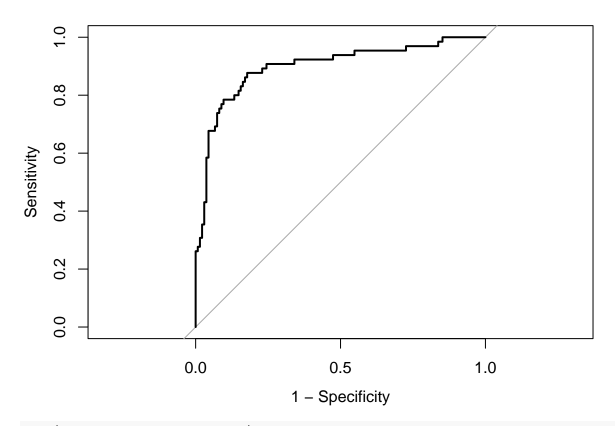
## Setting direction: controls < cases

roc.qda <- roc(y_test, qda.pred2[,2])

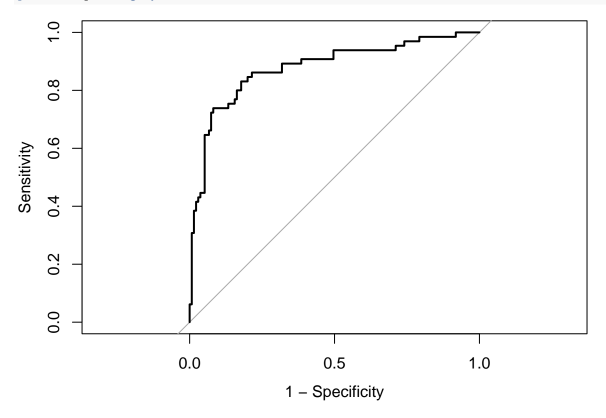
## Setting levels: control = 0, case = 1</pre>
```

## Setting direction: controls < cases

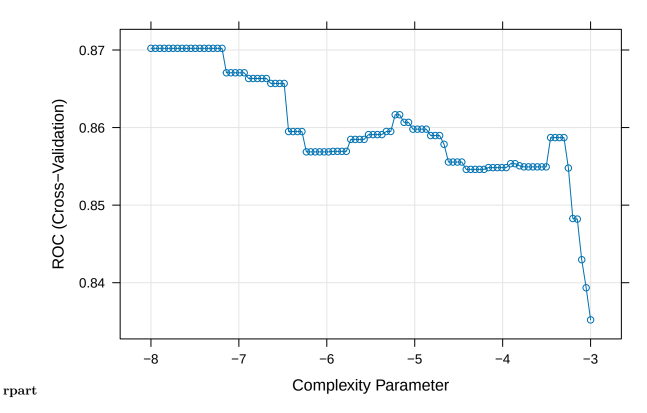
plot(roc.lda, legacy.axes = TRUE)



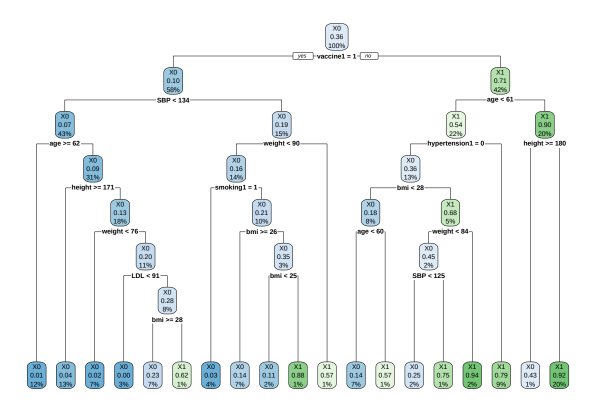
plot(roc.qda, legacy.axes = TRUE)

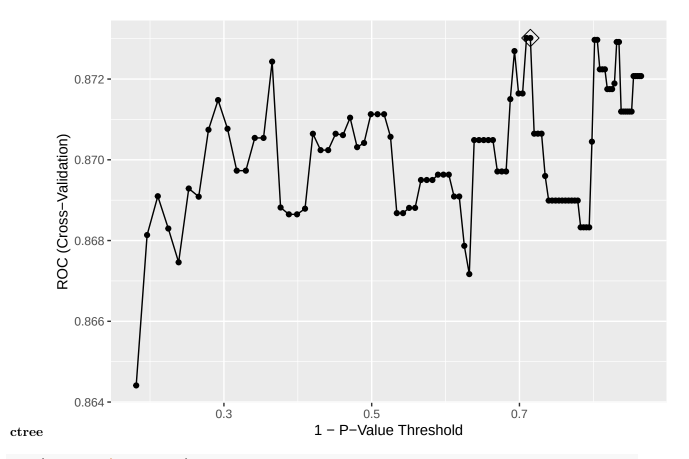


classification tree

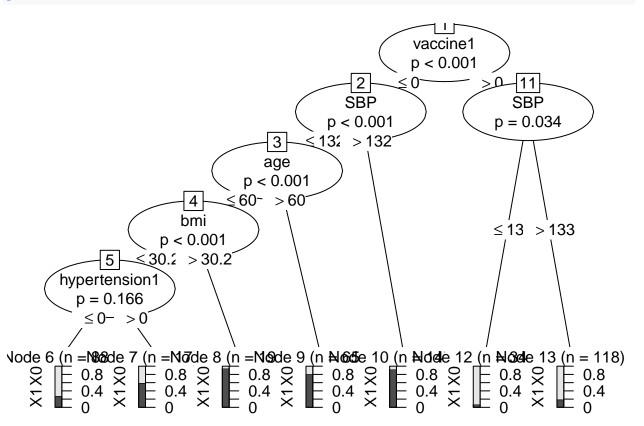


rpart.plot(rpart.fit\$finalModel)

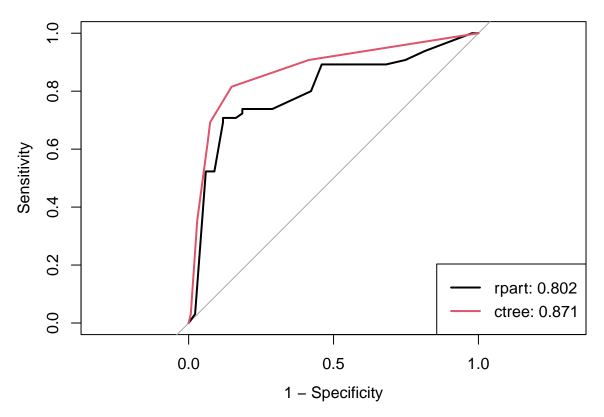








```
ctree.fit$bestTune
##
     mincriterion
## 59
        0.7147945
summary(resamples(list(rpart.fit, ctree.fit)))
classification tree test performance
##
## Call:
## summary.resamples(object = resamples(list(rpart.fit, ctree.fit)))
##
## Models: Model1, Model2
## Number of resamples: 10
##
## ROC
                      1st Qu.
                                 Median
                                                     3rd Qu.
               Min.
                                              Mean
## Model1 0.7990196 0.8544806 0.8699532 0.8702149 0.8843845 0.9240716
## Model2 0.7650560 0.8563843 0.8715717 0.8730165 0.8932019 0.9512599
##
## Sens
##
               Min.
                      1st Qu.
                                  Median
                                              Mean
                                                     3rd Qu.
## Model1 0.8039216 0.8486991 0.9038462 0.8909502 0.9411765 0.9607843
                                                                           0
## Model2 0.8039216 0.8829186 0.9125189 0.9045626 0.9366516 0.9803922
##
## Spec
##
               Min.
                      1st Qu.
                                 Median
                                              Mean
                                                     3rd Qu.
                                                                   Max. NA's
## Model1 0.4642857 0.6813424 0.7019704 0.7051724 0.7844828 0.8275862
## Model2 0.5357143 0.7142857 0.7586207 0.7399015 0.8189655 0.8620690
rpart.pred2 <- predict(rpart.fit, newdata = test.raw,</pre>
                       type = "prob")[,1]
ctree.pred <- predict(ctree.fit, newdata = test.raw, type = "prob")[,1]</pre>
roc.rpart <- roc(y_test, rpart.pred2)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls > cases
roc.ctree <- roc(y_test, ctree.pred)</pre>
## Setting levels: control = 0, case = 1
## Setting direction: controls > cases
```



#### random forest

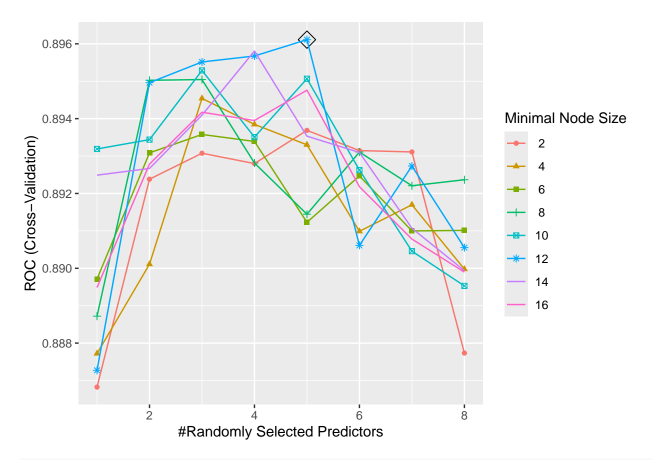
## ranger

```
## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not ## in the result set. ROC will be used instead.
```

## ggplot(rf.fit, highlight = TRUE)

## Warning: The shape palette can deal with a maximum of 6 discrete values because more
## than 6 becomes difficult to discriminate
## i you have requested 8 values. Consider specifying shapes manually if you need
## that many have them.

## Warning: Removed 16 rows containing missing values or values outside the scale range
## ('geom\_point()').



### rf.fit\$bestTune

```
## mtry splitrule min.node.size
## 38 5 gini 12
```

```
# test
rf.pred <- predict(rf.fit, newdata = test.raw, type = "prob")[,1]
roc.rf <- roc(test.raw$severity, rf.pred)</pre>
```

```
## Setting levels: control = 0, case = 1
## Setting direction: controls > cases
```

```
roc.rf

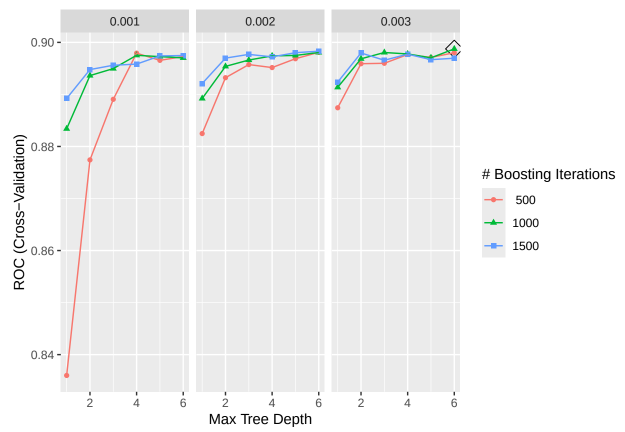
##

## Call:
## roc.default(response = test.raw$severity, predictor = rf.pred)
##

## Data: rf.pred in 135 controls (test.raw$severity 0) > 65 cases (test.raw$severity 1).
## Area under the curve: 0.8683

1-roc.rf$auc[1]

## [1] 0.1317379
```



### AdaBoost

```
# test
gbmA.pred <- predict(gbmA.fit, newdata = test.raw, type = "prob")[,1]
roc.gbmA <- roc(test.raw$severity, gbmA.pred)

## Setting levels: control = 0, case = 1

## Setting direction: controls > cases

1-roc.gbmA$auc[1]

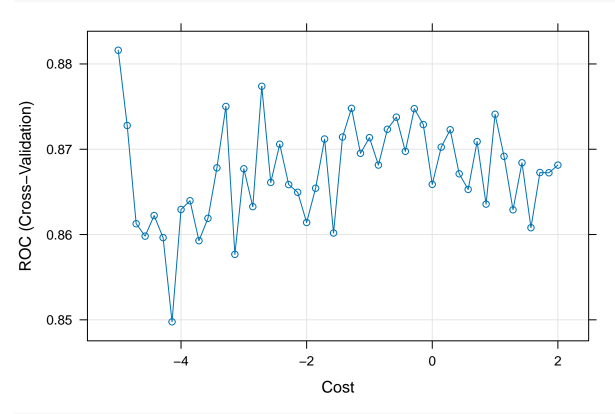
## [1] 0.1120228
```

 $\mathbf{svm}$ 

 $\operatorname{\mathbf{svml}}$ 

## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not ## in the result set. ROC will be used instead.

```
plot(svml.fit, highlight = TRUE, xTrans = log)
```

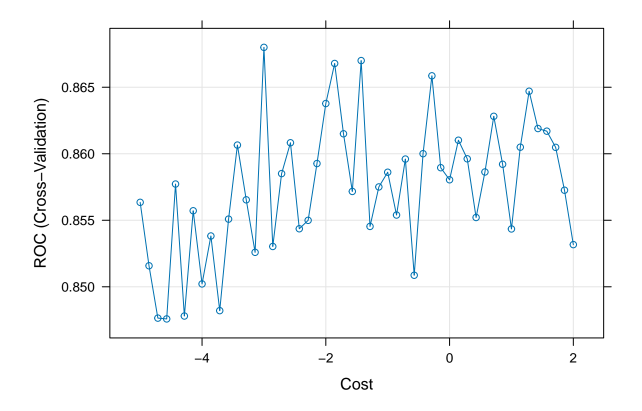


### svml.fit\$bestTune

```
## C
## 1 0.006737947
```

## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not ## in the result set. ROC will be used instead.

```
plot(svml.fit2, highlight = TRUE, xTrans = log)
```



### $\mathbf{svmr}$

## line search fails -0.6832979 0.3122083 1.048723e-05 -3.011969e-06 -3.814446e-08 1.354287e-08 -4.4082

# Sigma 0.000372699966223616 0.00305959206434424 0.00056783242423576 0.00466148574327131 0.000865129303016903 0.00710207402743375 0.00131808026275682 0.0108204676081991 0.00200818024890684 0.0164856799306543 0.90 ROC (Cross-Validation) 0.88 0.86 0.84 0.82 0.80 0 200 400 600 800 1000 Cost

#### svmr.fit\$bestTune

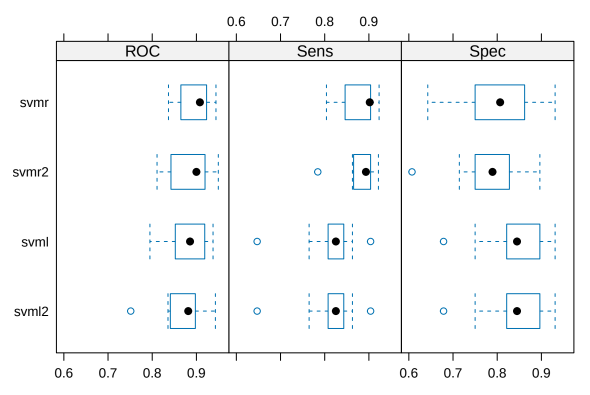
```
## sigma C
## 867 0.0005678324 526.0026
```

## model comparison

```
##
## Call:
## summary.resamples(object = resamp)
##
## Models: svmr, svmr2, svml, svml2
## Number of resamples: 10
```

```
##
## ROC
##
                     1st Qu.
                                Median
                                            Mean
                                                    3rd Qu.
        0.8368347 0.8719272 0.9079615 0.8982876 0.9211786 0.9442971
  svmr2 0.8109244 0.8503876 0.8999974 0.8863974 0.9173337 0.9496021
  svml 0.7948179 0.8566873 0.8856247 0.8816084 0.9149763 0.9376658
                                                                         0
   svml2 0.7514006 0.8419118 0.8815013 0.8679987 0.8968795 0.9429708
##
## Sens
##
                     1st Qu.
                                Median
                                                    3rd Qu.
              Min.
                                            Mean
## svmr 0.8039216 0.8552036 0.9019608 0.8812217 0.9033748 0.9230769
## svmr2 0.7843137 0.8696267 0.8932881 0.8812594 0.9033748 0.9215686
                                                                         0
   svml 0.6470588 0.8116516 0.8252262 0.8130090 0.8390837 0.9038462
                                                                         0
  svml2 0.6470588 0.8116516 0.8252262 0.8130090 0.8390837 0.9038462
##
## Spec
##
                     1st Qu.
                                Median
              Min.
                                            Mean
                                                    3rd Qu.
        0.6428571 0.7521552 0.8066502 0.8032020 0.8534483 0.9310345
## symr2 0.6071429 0.7521552 0.7894089 0.7788177 0.8189655 0.8965517
                                                                         0
## svml 0.6785714 0.8214286 0.8448276 0.8381773 0.8879310 0.9310345
                                                                         0
## syml2 0.6785714 0.8214286 0.8448276 0.8381773 0.8879310 0.9310345
```

### bwplot(resamp)



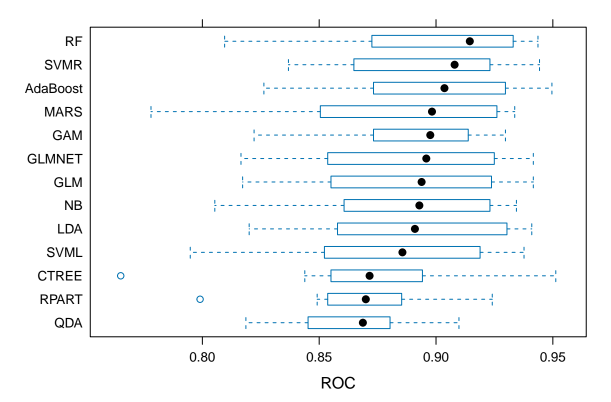
```
# test error
pred.svml <- predict(svml.fit, newdata = test.raw)
pred.svmr <- predict(svmr.fit, newdata = test.raw)
levels(pred.svml) <- levels(test.raw$severity)
confusionMatrix(data = pred.svml, reference = test.raw$severity)</pre>
```

```
## Confusion Matrix and Statistics
##
             Reference
##
              0 1
## Prediction
##
            0 115 12
            1 20 53
##
##
##
                  Accuracy: 0.84
##
                    95% CI: (0.7817, 0.8879)
##
       No Information Rate: 0.675
##
       P-Value [Acc > NIR] : 9.736e-08
##
##
                     Kappa: 0.6466
##
##
    Mcnemar's Test P-Value: 0.2159
##
##
               Sensitivity: 0.8519
##
               Specificity: 0.8154
##
            Pos Pred Value: 0.9055
##
            Neg Pred Value: 0.7260
##
                Prevalence: 0.6750
##
            Detection Rate: 0.5750
##
      Detection Prevalence: 0.6350
##
         Balanced Accuracy: 0.8336
##
##
          'Positive' Class: 0
##
levels(pred.svmr) <- levels(test_data$severity)</pre>
confusionMatrix(data = pred.svmr, reference = test.raw$severity)
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction 0
##
            0 120 16
            1 15 49
##
##
##
                  Accuracy: 0.845
                    95% CI: (0.7873, 0.8922)
##
##
       No Information Rate: 0.675
##
       P-Value [Acc > NIR] : 3.744e-08
##
##
                     Kappa: 0.6453
##
##
    Mcnemar's Test P-Value : 1
##
               Sensitivity: 0.8889
##
##
               Specificity: 0.7538
##
            Pos Pred Value: 0.8824
##
            Neg Pred Value: 0.7656
##
                Prevalence: 0.6750
##
            Detection Rate: 0.6000
##
      Detection Prevalence: 0.6800
```

```
## Balanced Accuracy : 0.8214
##

"Positive' Class : 0
##
```

## total model comparison



# Final model

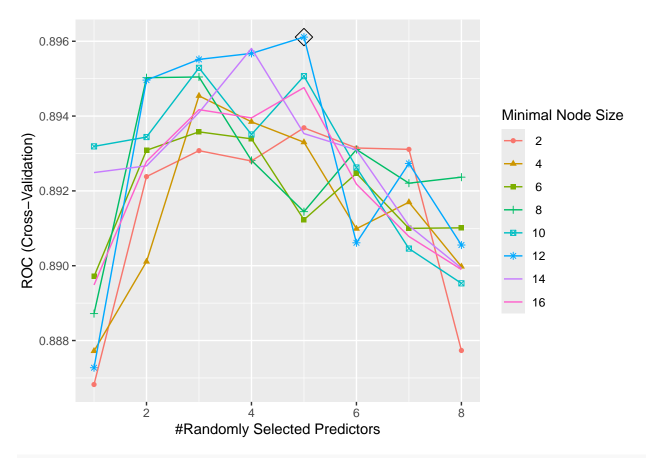
## $\mathbf{RF}$

```
ggplot(rf.fit, highlight = TRUE)
```

## Warning: The shape palette can deal with a maximum of 6 discrete values because more ## than 6 becomes difficult to discriminate

## i you have requested 8 values. Consider specifying shapes manually if you need
## that many have them.

## Warning: Removed 16 rows containing missing values or values outside the scale range
## ('geom\_point()').



## rf.fit\$bestTune

```
## mtry splitrule min.node.size
## 38 5 gini 12
```

```
# test
rf.pred <- predict(rf.fit, newdata = test.raw, type = "prob")[,1]
roc.rf <- roc(test.raw$severity, rf.pred)</pre>
```

## Setting levels: control = 0, case = 1

```
## Setting direction: controls > cases
```

### roc.rf

```
##
## Call:
## roc.default(response = test.raw$severity, predictor = rf.pred)
##
## Data: rf.pred in 135 controls (test.raw$severity 0) > 65 cases (test.raw$severity 1).
## Area under the curve: 0.8683
```

# 1-roc.rf\$auc[1]

## [1] 0.1317379

## $\mathbf{SVMR}$

```
# variable importance
svmr_importance <- varImp(svmr.fit, scale = TRUE)
plot(svmr_importance)</pre>
```

