# HW2-solution.R

### Heramb

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```
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#CS-422 HW-2
library (rpart)
library(caret)
## Warning: package 'caret' was built under R version 3.4.2
## Loading required package: lattice
## Loading required package: ggplot2
## Warning: package 'ggplot2' was built under R version 3.4.2
library (rpart.plot)
## Warning: package 'rpart.plot' was built under R version 3.4.2
library (ROCR)
## Warning: package 'ROCR' was built under R version 3.4.2
## Loading required package: gplots
## Warning: package 'gplots' was built under R version 3.4.2
##
## Attaching package: 'gplots'
## The following object is masked from 'package:stats':
##
##
      lowess
```

```
setwd("C:/Program Files/RStudio/Data FIles/")
rm(list=ls())

ilpd=read.csv("ILPD.csv", header = T, sep=",")

set.seed(100)
# Splitting into 60-40 (train-test).
index_ilpd<- sample(1:nrow(ilpd), size = 0.6*nrow(ilpd))
train_ilpd<- ilpd[index_ilpd, ]
test_ilpd<- ilpd[-index_ilpd, ]
# There are 234 and 349 instances to test and train the data

#a.For the training dataset, produce a correlation scatterplot of the variables.

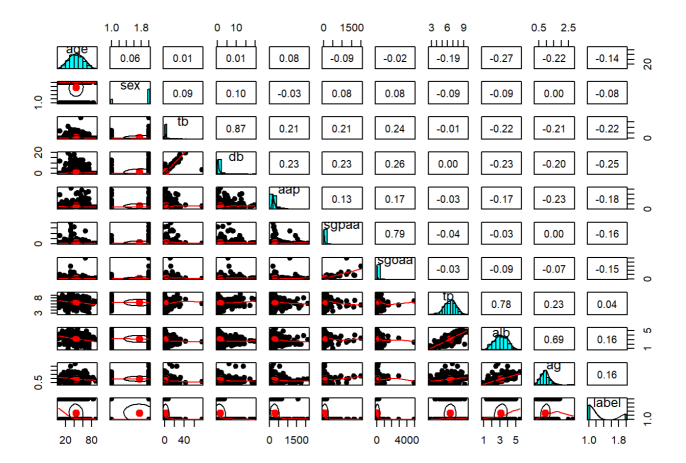
library(psych)</pre>
```

```
## Warning: package 'psych' was built under R version 3.4.2
```

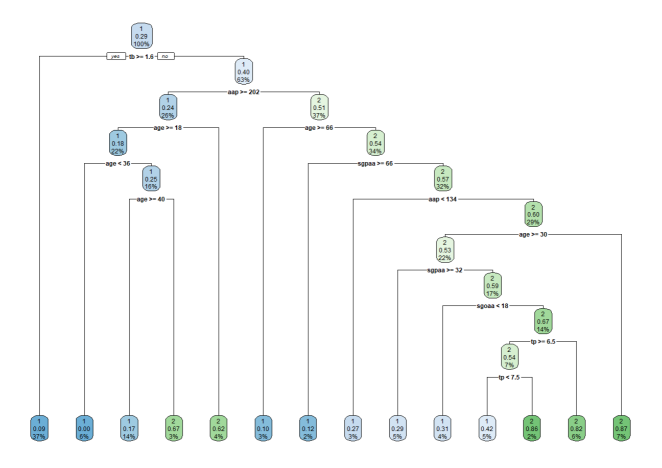
```
##
## Attaching package: 'psych'
```

```
## The following objects are masked from 'package:ggplot2':
##
## %+%, alpha
```

```
pairs.panels(ilpd, pch=19)
```



```
#i)Strongest correlated pair :- db(Direct Bilirubin) and tb(Total Bilirubin)
#ii) Weakest correlated pair :- tp(Total Proteins) and db(Direct Bilirubin), sex an
d ag(Ratio of Albumin to Globulin), sgpaa(Sgpt Alamine Aminotransferase) and ag(Rati
o of Albumin to Globulin)
#iii)Most negatively correlated :- age and alb(Albumin)
#iv) Variables appear to follow a Gaussian distribution :- age, tp(Total Proteins),
alb(Albumin), ag(Ratio of Albumin to Globulin)
#b)Yes, I think normalising or scaling the attributes will help the classification
task. Because, normalising the attributes will result in more linear relationship.
It will also help in providing a robust correlation. There is no point in normalisi
ng the data which is linear.
  #Attributes with non-linear range of values that should be normalised are: - Age,
tp(Total Proteins), alb(Albumin), ag(Ratio of Albumin to Globulin)
#C)
model <- rpart(label~ ., method = "class", data =train ilpd)</pre>
rpart.plot(model)
```



```
pred <- predict(model, test_ilpd, type = "class")
confusionMatrix(pred, test_ilpd[,11], positive = "1")</pre>
```

```
## Confusion Matrix and Statistics
##
##
          Reference
## Prediction 1 2
##
      1 145 51
          2 22 16
##
##
##
                Accuracy: 0.688
##
                  95% CI: (0.6244, 0.7468)
    No Information Rate: 0.7137
##
##
     P-Value [Acc > NIR] : 0.826721
##
##
                   Kappa : 0.123
## Mcnemar's Test P-Value : 0.001049
##
             Sensitivity: 0.8683
##
             Specificity: 0.2388
##
##
          Pos Pred Value : 0.7398
          Neg Pred Value : 0.4211
##
##
              Prevalence: 0.7137
##
          Detection Rate : 0.6197
## Detection Prevalence : 0.8376
##
      Balanced Accuracy: 0.5535
##
##
        'Positive' Class : 1
##
```

```
#Accuracy: 68.8%

#TPR(Sensitivity): 0.8683

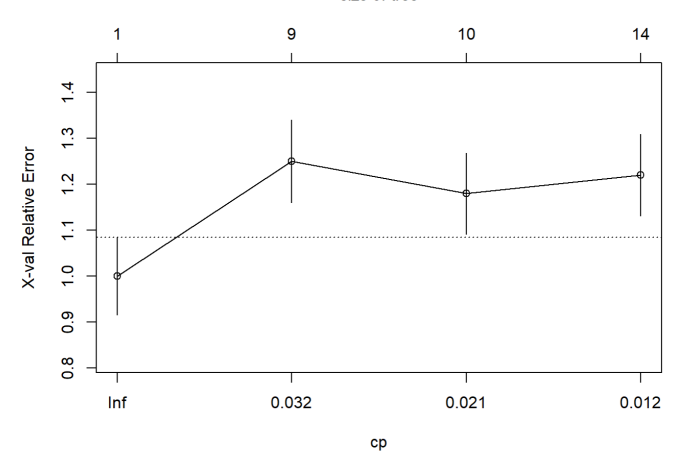
#TNR(Specificity): 0.2388

#PPV(Pos Pred Value): 0.7398

#d)
```

plotcp(model)

### size of tree



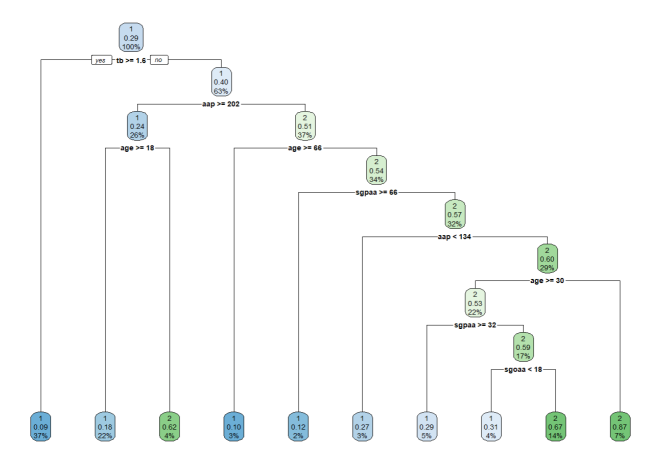
#### printcp(model)

```
##
## Classification tree:
## rpart(formula = label ~ ., data = train ilpd, method = "class")
##
## Variables actually used in tree construction:
## [1] aap age sgoaa sgpaa tb tp
##
## Root node error: 100/349 = 0.28653
##
## n= 349
##
          CP nsplit rel error xerror
## 1 0.033333
               0
                        1.00
                             1.00 0.084467
## 2 0.030000
                  8
                        0.67
                              1.25 0.089571
## 3 0.015000
                 9
                        0.64 1.18 0.088376
## 4 0.010000
               13
                        0.58
                             1.22 0.089080
```

```
model.pruned <- prune(model, cp = 0.021)
pred.pruned <- predict(model.pruned, test_ilpd, type = "class")
confusionMatrix(pred.pruned, test_ilpd[, 11], positive = "1")</pre>
```

```
## Confusion Matrix and Statistics
##
##
          Reference
## Prediction 1 2
      1 142 46
##
          2 25 21
##
##
##
                Accuracy: 0.6966
##
                  95% CI: (0.6333, 0.7548)
     No Information Rate : 0.7137
##
##
     P-Value [Acc > NIR] : 0.74428
##
##
                   Kappa : 0.1807
## Mcnemar's Test P-Value : 0.01762
##
             Sensitivity: 0.8503
##
             Specificity: 0.3134
##
##
          Pos Pred Value : 0.7553
          Neg Pred Value : 0.4565
##
##
              Prevalence: 0.7137
           Detection Rate: 0.6068
##
## Detection Prevalence : 0.8034
##
       Balanced Accuracy: 0.5819
##
##
        'Positive' Class : 1
##
```

```
rpart.plot(model.pruned)
```

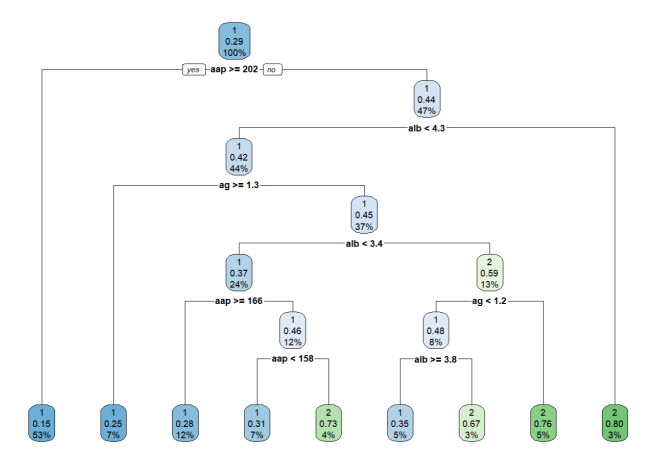


#Accuracy of the model is 68.8% and by changing the values of cp I got a better accuracy of 69.66%.

#I think that with better accuracy after pruning it reduces the size of learning tr ee. Thus, because of reduced cost of complexity it increases the accuracy.

```
#e)Build a model
```

newmodel<- rpart(label ~ alb+ag+aap, method = "class", data = train\_ilpd)
rpart.plot(newmodel)</pre>



```
newpred <- predict(newmodel,test_ilpd,type = "class")
confusionMatrix(newpred,test_ilpd[,11], positive = "1")</pre>
```

```
## Confusion Matrix and Statistics
##
##
          Reference
## Prediction 1 2
##
       1 152 49
          2 15 18
##
##
##
                Accuracy: 0.7265
                  95% CI: (0.6646, 0.7825)
##
##
    No Information Rate: 0.7137
##
     P-Value [Acc > NIR] : 0.3623
##
##
                   Kappa : 0.2109
## Mcnemar's Test P-Value: 3.707e-05
##
##
             Sensitivity: 0.9102
##
             Specificity: 0.2687
##
          Pos Pred Value: 0.7562
          Neg Pred Value : 0.5455
##
##
             Prevalence: 0.7137
          Detection Rate: 0.6496
##
##
    Detection Prevalence: 0.8590
      Balanced Accuracy: 0.5894
##
##
##
        'Positive' Class : 1
##
```

#### summary(ilpd)

```
## age
                              tb
                                            db
                  sex
## Min. : 4.00 Female:142 Min. : 0.400 Min. : 0.100
## 1st Qu.:33.00 Male :441 1st Qu.: 0.800 1st Qu.: 0.200
## Median :45.00
                         Median: 1.000 Median: 0.300
## Mean :44.75
                          Mean : 3.299 Mean : 1.486
## 3rd Qu.:58.00
                         3rd Qu.: 2.600 3rd Qu.: 1.300
                          Max. :75.000 Max. :19.700
## Max. :90.00
## aap
                   sqpaa
                                 sgoaa
                                                tp
## Min. : 63.0 Min. : 10.00 Min. : 10.0 Min. :2.700
## 1st Qu.: 175.5 1st Qu.: 23.00 1st Qu.: 25.0 1st Qu.:5.800
## Median: 208.0 Median: 35.00 Median: 42.0 Median: 6.600
## Mean : 290.6 Mean : 80.71 Mean : 109.9 Mean :6.483
## 3rd Qu.: 298.0 3rd Qu.: 60.50 3rd Qu.: 87.0 3rd Qu.:7.200
## Max. :2110.0 Max. :2000.00 Max. :4929.0 Max. :9.600
                   ag
## alb
                               label
## Min. :0.900 Min. :0.300 Min. :1.000
## 1st Qu.:2.600 1st Qu.:0.700 1st Qu.:1.000
## Median :3.100 Median :0.940 Median :1.000
## Mean :3.142 Mean :0.947 Mean :1.286
## 3rd Qu.:3.800 3rd Qu.:1.100 3rd Qu.:2.000
## Max. :5.500 Max. :2.800 Max. :2.000
```

```
## [1] 16.18983
sd(ilpd$sex)
\#\# Warning in var(if (is.vector(x) || is.factor(x)) x else as.double(x), na.rm = na
.rm): Calling var(x) on a factor x is deprecated and will become an error.
\#\# Use something like 'all(duplicated(x)[-1L])' to test for a constant vector.
## [1] 0.4296034
sd(ilpd$tb)
## [1] 6.209522
sd(ilpd$db)
## [1] 2.808498
sd(ilpd$aap)
## [1] 242.938
sd(ilpd$sgpaa)
## [1] 182.6204
sd(ilpd$sgoaa)
## [1] 288.9185
sd(ilpd$tp)
## [1] 1.085451
sd(ilpd$alb)
## [1] 0.7955188
sd(ilpd$ag)
## [1] 0.3184925
```

```
# Accuracy :- 72.65%(Increased by 3.85%)
#TPR(Sensitivity) : 0.9102(Increased by 4.19%)
#TNR(Specificity) : 0.2687(Increased by 2.99%)
#PPV(Pos Pred Value) : 0.7562(Increased by 1.64%)

#f)
#(i) a ROC curve using the ROCR package.
#ROC for model
pred.roc <- predict(model,newdata=test_ilpd,type="prob")[,2]
f.pred <- prediction(pred.roc,test_ilpd$label)
f.perf <- performance(f.pred, "tpr", "fpr")
plot(f.perf, colorize=T, lwd=3, main = "ROC")
abline(0,1)</pre>
```

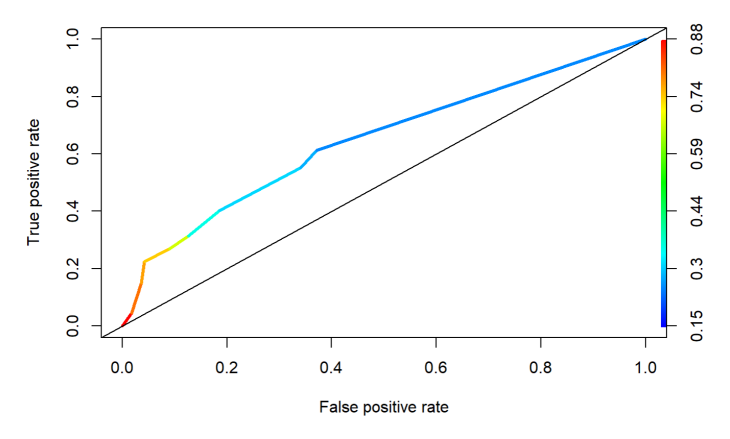
# 

```
auc<-performance(f.pred, measure = "auc")
auc@y.values[[1]]</pre>
```

```
## [1] 0.6675753
```

```
#ROC for newmode1
pred.roc1 <- predict(newmodel,newdata=test_ilpd,type="prob")[,2]
f.pred1 <- prediction(pred.roc1,test_ilpd$label)
f.perf1 <- performance(f.pred1, "tpr", "fpr")
plot(f.perf1, colorize=T, lwd=3, main = "ROC")
abline(0,1)</pre>
```

## **ROC**



```
auc<-performance(f.pred1, measure = "auc")
auc@y.values[[1]]</pre>
```

```
## [1] 0.6455
```

```
#ii) AUC for model: 66.75%
# AUC for newmodel: 64.55%

#iii) Previous Model is better because the auc is closest to 1 as compared to newmodel.
```