MVA Assignment 8

Assignment 8 - Logistic regression

This document performs Logistic Regression on the Heart Failure Prediction dataset. We iterate over multiple models to come up with the most robust model.

Let us load libraries and data

```
# clear environment rm(list = ls())
# defining libraries
library(ggplot2) library(dplyr)
library(PerformanceAnalytics)
library(data.table) library(sqldf)
library(nortest) library(MASS)
library(rpart) library(class)
library(ISLR) library(scales)
library(ClustOfVar) library(GGally)
library(reticulate) library(ggthemes)
library(RColorBrewer)
library(gridExtra) library(kableExtra)
library(Hmisc) library(corrplot)
library(energy) library(nnet)
library(Hotelling) library(car)
library(devtools) library(ggbiplot)
```

reading data

library(factoextra)

library(rgl) library(FactoMineR) library(psych) library(nFactors) library(scatterplot3d) library(Imtest)

library(mctest) library(aod) library(InformationValue) library(pROC) library(tidyverse) library(caret) library(Information)

data <- read.csv('/Users/mac/Downloads/heart_failure_clinical records dataset.csv') str(data)

'data.frame': 299 obs. of 13 variables:

\$ age : num 75 55 65 50 65 90 75 60 65 80 ...

\$ anaemia : int 0 0 0 1 1 1 1 1 0 1 ...

##\$ creatinine_phosphokinase: int 582 7861 146 111 160 47 246 315 157 123 ...

\$ diabetes : int 0 0 0 0 1 0 0 1 0 0 ... ## \$ ejection fraction : int 20 38 20 20 20 40 15 60 65 35 ...

##\$high_blood_pressure : int 1000010001...

##\$ platelets : num 265000 263358 162000 210000 327000 ... ##\$ serum_creatinine : num 1.9 1.1 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.4 ... ##\$ serum sodium : int 130 136129137116132 137 131138133 ...

##\$ sex : int 1 1 1 1 0 1 1 1 0 1 ... ##\$ smoking : int 0 0 1 0 0 1 0 1 0 1 ... ##\$ time : int 4 6 7 7 8 8 10 10 10 10 ... ##\$ DEATH EVENT : int 1 1 1 1 1 1 1 1 ...

Fitting a logistic regression model

We recall three key points from third assignment (EDA) -

- 1. Our data has no missing values
- 2. We saw 4 observations as outliers
- 3. We saw no multicollinearity as our VIF values were all below 2 Hence we have a very small

pre-processing step of removing outliers.

Data Cleaning - Let's remove these outliers

```
data <- data[data$ejection_fraction <70,] data <-
data[data$creatinine_phosphokinase <7000,]
str(data)
```

'data.frame': 295 obs. of 13 variables:

\$ age : num 75 65 50 65 90 75 60 65 80 75 ...

\$ anaemia : int 0 0 1 1 1 1 1 0 1 1 ...

##\$ creatinine_phosphokinase: int 582 146 111 160 47 246 315 157 123 81 ...

\$ diabetes : int 0 0 0 1 0 0 1 0 0 0 ... ## \$ ejection_fraction : int 20 20 20 20 40 15 60 65 35 38 ...

##\$high_blood_pressure : int 1000100011...

##\$ platelets : num 265000 162000 210000 327000 204000 ... ##\$ serum_creatinine : num 1.9 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.44 ... ##\$ serum_sodium : int 130 129 137 116 132 137 131 138 133 131 ...

##\$ sex : int 1 1 1 0 1 1 1 0 1 1 ... ##\$ smoking : int 0 1 0 0 1 0 1 0 1 1 ...

```
## $ time
                                     : int 4 7 7 8 8 10 10 10 10 10 ...
                                     : int 1111111111...
## $ DEATH_EVENT
We remove the 4 outliers before proceeding to modeling exercise. Converting
categorical features and dependent variable to factor
data$DEATH EVENT <- factor(data$DEATH EVENT)
data$anaemia <- factor(data$anaemia) data$diabetes <-
factor(data$diabetes)
data$high blood pressure <- factor(data$high blood pressure)data$sex <-
factor(data$sex) data$smoking <- factor(data$smoking) str(data)</pre>
## 'data.frame':
                             295 obs. of 13 variables:
## $ age
                                           : num 75 65 50 65 90 75 60 65 80 75 ...
                                               : Factor w/ 2 levels "0", "1": 1 1 2 2 2 2 2 1 2 2 ...
## $ anaemia
##$ creatinine_phosphokinase: int 582 146 111 160 47 246 315 157 123 81 ...
## $ diabetes
                                               : Factor w/ 2 levels "0","1": 1 1 1 2 1 1 2 1 1 1 ...
## $ ejection fraction
                                            : int 20 20 20 20 40 15 60 65 35 38 ...
## $ high_blood_pressure
                                               : Factor w/ 2 levels "0", "1": 2 1 1 1 2 1 1 1 2 2 ...
                                          : num 265000 162000 210000 327000 204000 ...
## $ platelets
                                              : num 1.9 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.44 ...
## $ serum creatinine
## $ serum sodium
                                            : int 130 129 137 116 132 137 131 138 133 131 ...
## $ sex
                                               : Factor w/ 2 levels "0", "1": 2 2 2 1 2 2 2 1 2 2 ...
## $ smoking
                                               : Factor w/ 2 levels "0", "1": 1211212121...
## $ time
                                            : int 477881010101010...
## $ DEATH EVENT
                                               : Factor w/ 2 levels "0","1": 2 2 2 2 2 2 2 2 2 2 ...
Two-way contingency table of categorical outcome and predictors Since we want to
make sure there are not 0 cells
xtabs(~DEATH EVENT + anaemia, data = data)
                 anaemia
## DEATH EVENT
        0 119 83 ##
                          148
45 xtabs(~DEATH EVENT+
diabetes, data = data)
##
                  diabetes
```

xtabs(~DEATH_EVENT + high_blood_pressure, data = data)

154

0 1

DEATH EVENT

0 117 85 ##

##

39

```
##
                high blood pressure
                  0 1
## DEATH_EVENT
##
               013666
               15637
xtabs(~DEATH EVENT + sex, data = data)
## DEATH EVENT
##
       070132##
                     133
60
xtabs(~DEATH_EVENT + smoking, data = data)
##
               smoking
## DEATH_EVENT
                 0
               013666
               16330
##
```

We note no 0 or low cells in any of the categorical variables.

Checking event rate in data - this will help determine prob. value for thresholds

```
##
## 0 1
## 202 93
We note a 31.5% event rate in the data.
```

Fitting a model

Iteration - 1 All variables

```
##
         data = data)
##
## Deviance Residuals:
                                       Max ## -2.1947 -
       Min
               10
                       Median 3Q
## Coefficients:
                                       Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                   1.127e+01 5.669e+00
                                                              1.988 0.04682 *
                                    5.197e-02 1.623e-02
                                                              3.202 0.00137 **
## age
                                     -8.307e-023.635e-01-0.2290.81923
## anaemia1
## creatinine_phosphokinase 1.101e-04 2.207e-04
                                                             0.499 0.61779
## diabetes1
                                    1.623e-01 3.528e-01
                                                             0.460 0.64551
## ejection fraction
                                -8.244e-021.723e-02-4.7841.72e-06***
## high_blood_pressure1
                                -2.472e-013.740e-01-0.6610.50859
## platelets
                                -1.069e-061.908e-06-0.5600.57538
## serum creatinine
                                4.669e-01 2.049e-01
                                                              2.279 0.02265 *
## serum sodium
                                -7.236e-023.970e-02-1.8230.06831.
## sex1
                                -5.384e-014.136e-01-1.3020.19294
## smoking1
                                -4.051e-024.196e-01-0.0970.92308
## time
                                -2.160e-023.114e-03-6.9364.04e-12***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
## (Dispersion parameter for binomial family taken to be 1) ##
       Null deviance: 367.71 on 294 degrees of freedom ## Residual
deviance: 215.95 on 282 degrees of freedom
## AIC: 241.95
## Number of Fisher Scoring iterations: 6
```

Key observations

- 1. We note age, ejection_fraction, serum_creatinine and time as significant variables in this iteration 2. The above iteration has an AIC of 241.95
- 3. Interpreting the coefficient For every one unit change in age, the log odds of death (versus survival) increases by 5.197e-02
- 4. None of the categorical variables are significant in predicting the death event

Predicting the outcome

```
glm.probs <- predict(mylogit,type = "response") glm.probs[1:5]

## 1 3 4 5 6

## 0.9802885 0.9671714 0.9008819 0.9956762 0.9417182
```

The first five probabilities in this case are all close to 1 as evidenced in the data as well.

Let's use a base calculation to figure out accuracy

```
# Here we try the case of using default 0.5 as threshold glm.pred <-
ifelse(glm.probs > 0.5, "1", "0") table(data$DEATH_EVENT,glm.pred)
```

```
## glm.pred
## 0 1
## 018517
## 12766
```

Looking at the diagonal, we're not that bad. We have an overall accuracy of ~85% (Diagonals summed over overall sum) But note this is called a base model because we didn't do splitting into train and test so the model trained on the entire data and predicted on the entire data. Without out of sample testing one cannot claim robustness as there may be overfitting here.

Deciding on optimal cutoff

```
optCutOff <- optimalCutoff(data$DEATH_EVENT, glm.probs)[1] optCutOff
```

##[1]0.4669198

We used 0.5 above to classify however we want the cut-off where model is balanced in accuracy measures. We note this as 0.466. This tells us that we can use this prob cutoff to classify an observation as 0 or 1. If the prob-value is below this threshold we can classify it as survival else death event.

Mis-classification error

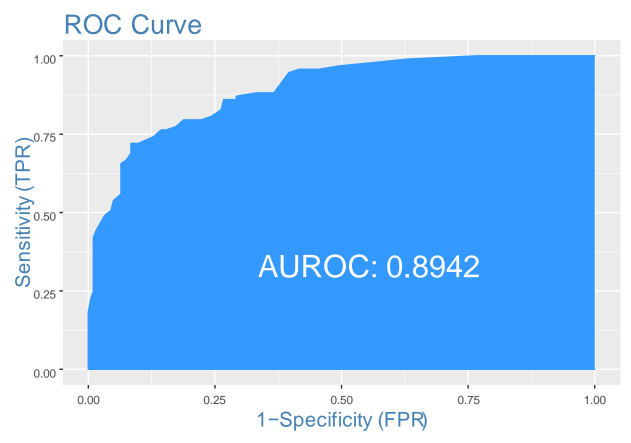
misClassError(data\$DEATH_EVENT, glm.probs, threshold = optCutOff)

##[1]0.1458

We note a misclassification error of 14.58%

ROC curve

plotROC(data\$DEATH_EVENT, glm.probs)

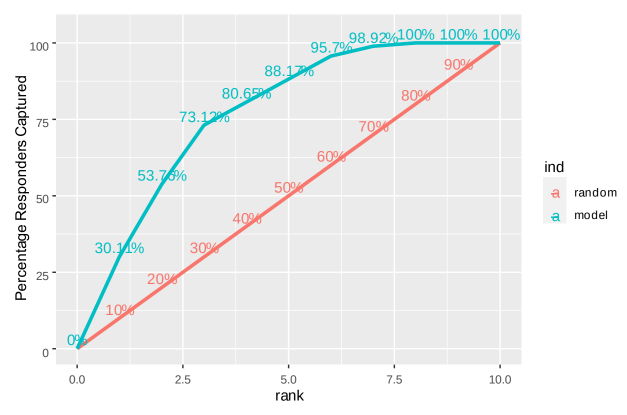


We see an AUC of 0.894 which is decent.

KS plot

ks_stat(data\$DEATH_EVENT, glm.probs)
[1] 0.6223
ks_plot(data\$DEATH_EVENT, glm.probs)

KS Plot



A KS plot answers the question how many responders/ deaths can we capture if we target x% of the population. Here, we see we can capture 73% responders if we target first 30% of the population.

Confusion Matrix and all accuracy measures

```
confusionMatrix(data = as.factor(glm.pred), reference = as.factor(data$DEATH_EVENT),
         mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction
                        1
        0 185 27 ##
##
                         1
1766
##
##
                       Accuracy: 0.8508
                           95% CI: (0.805, 0.8895)
##
           No Information Rate: 0.6847
##
##
            P-Value [Acc > NIR]: 4.394e-11
##
##
                          Kappa: 0.6442
##
## Mcnemar's Test P-Value: 0.1748##
##
        Precision: 0.8726##
                                 Recall:
0.9158
```

```
##
##
##
##
##
F1:0.8937
Prevalence:0.6847
Detection Rate:0.6271
Detection Prevalence:0.7186##
Balanced Accuracy:0.8128
##
##
"Positive' Class:0
```

We have to be careful here. From above, we can see accuracy of the model is 85.0% as before. Precision is 0.87 and Recall is 0.91 while F1- score is 0.89 but this is for positive class taken as '0'. However, we want to understand precision, recall for positive class as predicting death events is more important than survival events.

Confusion Matrix and all accuracy measures for positive class chosen as death=1

```
confusionMatrix(data = as.factor(glm.pred), reference = as.factor(data$DEATH_EVENT),positive='1',
    mode = "prec recall")
## Confusion Matrix and Statistics
##
                Reference
##
## Prediction
                   0
                        1
        0 185 27 ##
##
1766
##
##
                       Accuracy: 0.8508
##
                           95% CI: (0.805, 0.8895)
##
           No Information Rate: 0.6847
##
            P-Value [Acc > NIR]: 4.394e-11
##
##
                          Kappa: 0.6442
##
## Mcnemar's Test P-Value: 0.1748##
        Precision: 0.7952##
##
                                Recall:
0.7097
##
                              F1:0.7500
##
        Prevalence: 0.3153 ##
                                Detection
Rate: 0.2237
##
        Detection Prevalence: 0.2814##
        Balanced Accuracy: 0.8128
##
##
               'Positive' Class: 1
##
```

We give positive class as '1' as we want to understand precision and recall of death events more than survival events so we know what to maximize for. Our accuracy of the model is 85.0% as before however precision is 0.79 and recall is 0.70 while F1- score is 0.75. This is a more true picture of our model than before and we know that the overall accuracy is slightly dominant towards predicting survival events better than death events.

Using optimal cutoff to determine accuracy measures

```
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit,type="response")>threshold,1,0) actual_values<-data$DEATH_EVENT
confusionMatrix(data = as.factor(predicted_values),
reference = as.factor(actual_values), mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction
        0 185 26 ##
                         1
##
1767
##
##
                       Accuracy: 0.8542
                           95% CI: (0.8087, 0.8925)
##
##
           No Information Rate: 0.6847
##
            P-Value [Acc > NIR]: 1.641e-11
##
##
                          Kappa: 0.6533
##
## Mcnemar's Test P-Value: 0.2225##
        Precision: 0.8768##
##
                                 Recall:
0.9158
##
                              F1: 0.8959
##
        Prevalence: 0.6847 ##
                                 Detection
Rate: 0.6271
##
        Detection Prevalence: 0.7153##
        Balanced Accuracy: 0.8181
##
                'Positive' Class: 0
##
```

Our new accuracy has gone up from 85% to 85.4% all by optimizing the prob. cutoff threshold.

Using optimal cutoff to determine accuracy measures with positive class as 1

```
##
##
##
##
                    Accuracy: 0.8542
                      95% CI: (0.8087, 0.8925)
        No Information Rate: 0.6847
        P-Value [Acc > NIR]: 1.641e-11
##
##
                         Kappa: 0.6533
##
## Mcnemar's Test P-Value: 0.2225##
##
       Precision: 0.7976##
                               Recall:
0.7204
##
                            F1:0.7571
       Prevalence: 0.3153 ## Detection
##
Rate: 0.2271
##
       Detection Prevalence: 0.2847##
       Balanced Accuracy: 0.8181
##
```

'Positive' Class: 1

##

##

Here, we see that we have improved our recall while keeping precision same and hence consequently our F1 score.

However we may be over-fitting here as we haven't kept a hold out set. This is something we will explore in future iterations.

Iteration - 2 Using only significant variables from Iteration 1

```
# Model 2 mylogit_2 <- glm(DEATH_EVENT
~ age
    +ejection fraction+
    serum_creatinine+time, data = data, family = "binomial")
summary(mylogit 2)
##
## Call:
## glm(formula = DEATH EVENT ~ age + ejection fraction + serum creatinine +
              time, family = "binomial", data = data)
##
## Deviance Residuals:
        Min
                1Q
                        Median 3Q
                                         Max ## -2.1419 -
##
## Coefficients:
##
        Estimate Std. Error z value Pr(>|z|) ## (Intercept)
                                                         0.61105
        1.069480.571 0.56776
## age
                          0.04813
                                        0.01533
                                                     3.140 0.00169 **
## ejection fraction -0.07854
                                          0.01628-4.8241.41e-06 ***
## serum creatinine
                                        0.19991
                                                     2.850 0.00438 * *
                          0.56967
                                          0.00292-7.0471.83e-12***
## time
                         -0.02058
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
## (Dispersion parameter for binomial family taken to be 1) ##
        Null deviance: 367.71 on 294 degrees of freedom ## Residual
deviance: 222.40 on 290 degrees of freedom
## AIC: 232.4
##
## Number of Fisher Scoring iterations: 5
 We note the lower AIC value in this iteration of 232.4
```

Predicting the outcome

```
glm.probs_2<-predict(mylogit_2,type="response") glm.probs_2[1:5]

## 1 3 4 5 6

## 0.97467640.9407823 0.9156974 0.9718717 0.9444430
```

The first five probabilities in this case are all close to 1 as evidenced in the data as well.

Let's use a base calculation to figure out accuracy

```
# Here we try the case of using default 0.5 as threshold glm.pred_2 <- ifelse(glm.probs_2 < 0.5, "0", "1")
```

```
##
##
##
table(data$DEATH_EVENT,glm.pred_2)

glm.pred_2
0 1
0 182 20 1
30 63
```

Looking at the diagonal, we have an overall accuracy of \sim 83% (Diagonals summed over overall sum). This is lower than before which makes sense since we removed the unnecessary independent variables But how do we know this is more robust than the model before ?

Deciding on optimal cutoff

```
optCutOff<-optimalCutoff(data$DEATH_EVENT, glm.probs_2)[1] optCutOff
## [1] 0.5989542
```

Mis-classification error

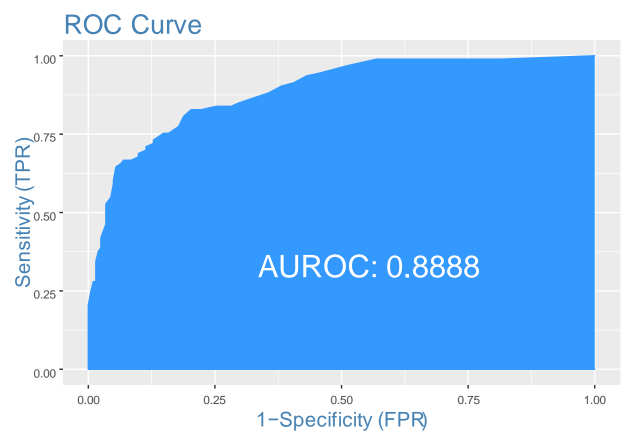
```
misClassError(data$DEATH_EVENT, glm.probs_2, threshold = optCutOff)
```

##[1]0.1492

We note a misclassification error of 14.92%

ROC curve

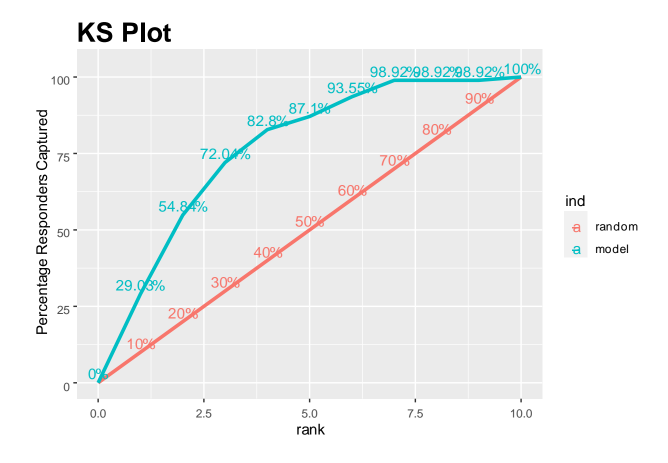
```
plotROC(data$DEATH_EVENT, glm.probs_2)
```



We see an AUC of 0.888 which is decent.

KS plot

```
ks_stat(data$DEATH_EVENT, glm.probs_2)
## [1] 0.6151
ks_plot(data$DEATH_EVENT, glm.probs_2)
```



Confusion Matrix and all accuracy measures

```
confusionMatrix(data = as.factor(glm.pred_2),
                                             reference = as.factor(data$DEATH_EVENT), mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction
                        1
        0 182 30 ##
                         1
##
2063
##
##
                       Accuracy: 0.8305
##
                           95% CI: (0.7827, 0.8715)
           No Information Rate: 0.6847
##
            P-Value [Acc > NIR]: 9.527e-09
##
##
                          Kappa: 0.5957
##
##
## Mcnemar's Test P-Value: 0.2031##
##
        Precision: 0.8585##
                                Recall:
0.9010
##
                              F1:0.8792
```

```
## Prevalence: 0.6847 ## Detection
Rate: 0.6169 ## Detection Prevalence: 0.7186
## Balanced Accuracy: 0.7892
##
## 'Positive' Class: 0
##
```

Our new accuracy is 83.0%. This is slightly lower than before which makes sense since we have eliminated some independent variables in this iteration.

Confusion Matrix and all accuracy measures for positive class chosen as death=1

```
confusionMatrix(data = as.factor(glm.pred_2), reference =
    as.factor(data$DEATH_EVENT),positive='1', mode = "prec_recall")
## Confusion Matrix and Statistics
##
                Reference
## Prediction
##
        0 182 30 ##
                        1
2063
##
##
                       Accuracy: 0.8305
                           95% CI: (0.7827, 0.8715)
##
##
           No Information Rate: 0.6847
##
            P-Value [Acc > NIR]: 9.527e-09
##
##
                          Kappa: 0.5957
##
## Mcnemar's Test P-Value: 0.2031##
##
        Precision: 0.7590##
                                Recall:
0.6774
                              F1:0.7159
##
##
        Prevalence: 0.3153 ## Detection
Rate: 0.2136
        Detection Prevalence: 0.2814##
##
        Balanced Accuracy: 0.7892
##
               'Positive' Class: 1
##
##
```

Our accuracy of the model is 83.0% as before however precision is 0.75 and recall is 0.67 while F1-score is 0.71. This is a more true picture of our model than before and we know that the overall accuracy is slightly dominant towards predicting survival events better than death events. We can see that all precision, recall, F1 score, accuracy and AUC are slightly lower in this iteration.

Using optimal cutoff to determine accuracy measures

```
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit_2,type="response")>threshold,1,0) actual_values<-datasdering DEATH_EVENT
confusionMatrix(data = as.factor(predicted_values), reference = as.factor(actual_values),
              mode = "prec_recall")
## Confusion Matrix and Statistics
##
                Reference
##
## Prediction
##
        0 191 33 ##
                         1
1160
##
##
                        Accuracy: 0.8508
##
                           95% CI: (0.805, 0.8895)
##
           No Information Rate: 0.6847
            P-Value [Acc > NIR]: 4.394e-11
##
##
##
                           Kappa: 0.631
##
## Mcnemar's Test P-Value: 0.001546
##
        Precision: 0.8527##
##
                                 Recall:
0.9455
                              F1:0.8967
##
##
        Prevalence: 0.6847 ##
                                 Detection
Rate: 0.6475
        Detection Prevalence: 0.7593##
##
        Balanced Accuracy: 0.7954
##
                'Positive' Class: 0
##
##
On using the optimal cutoff however, Our new accuracy has gone up from 83% to 85.0% all by optimizing the prob.
cutoff threshold.
Using optimal cutoff to determine accuracy measures with positive class as 1
threshold=optCutOff
predicted values<-ifelse(predict(mylogit 2,type="response")>threshold,1,0) actual values<-data$DEATH EVENT
confusionMatrix(data = as.factor(predicted values), reference =
                   as.factor(actual values), positive='1', mode =
                   "prec_recall")
## Confusion Matrix and Statistics
##
                Reference
##
## Prediction
                        1
        0 191 33 ##
##
                         1
1160
```

##

Accuracy: 0.8508

95% CI: (0.805, 0.8895)

No Information Rate : 0.6847 ## P-Value [Acc> NIR] : 4.394e-11

##

Kappa: 0.631

##

Mcnemar's Test P-Value: 0.001546

##

Precision: 0.8451## Recall:

0.6452

F1:0.7317 ## Prevalence:0.3153 ## Detection

Rate: 0.2034

Detection Prevalence : 0.2407 ## Balanced Accuracy : 0.7954

##

'Positive' Class: 1

##

Here, we see that we have improved our pecision however recall is much worse.

The problem in the first two iterations however is that we haven't kept a test set and may have over-fitted the model unknowingly.

Iteration - 3 All variables but splitting into train and test

Splitting into train and test - 70%, 30% split

We see our train data has event rate of 31.7% and our test data has event rate of 31.0%. This can happen in reality as well and hence a good accuracy on test will ensure we have built a robust model.

We will now train the model on training set and test on test set.

```
# Model 3
mylogit_3 <- glm(DEATH_EVENT ~ age+anaemia+creatinine_phosphokinase+
    diabetes+ejection fraction+high blood pressure+platelets+
    serum_creatinine+serum_sodium+sex+smoking+time,
    data = train_data, family = "binomial")
summary(mylogit 3)
##
## Call:
## glm(formula = DEATH_EVENT~ age + anaemia + creatinine_phosphokinase +
                diabetes + ejection fraction + high blood pressure + platelets +
##
##
                serum creatinine + serum sodium + sex + smoking + time, family = "binomial",
##
          data = train_data)
##
## Deviance Residuals:
                                       Max ## -2.0286 -
##
       Min
               1Q
                       Median 3Q
##
## Coefficients:
##
                                       Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                   1.240e+016.840e+00
                                                              1.813 0.06989.
## age
                                    5.037e-02 1.910e-02
                                                              2.637 0.00835 **
## anaemia1
                                                             0.865 0.38686
                                    3.813e-01 4.407e-01
## creatinine phosphokinase 6.435e-05 2.707e-04
                                                             0.238 0.81212
## diabetes1
                                    2.028e-01 4.338e-01
                                                             0.467 0.64017
## ejection_fraction
                                     -1.024e-012.208e-02-4.6363.55e-06***
```

```
## high blood pressure1 -4.740e-014.734e-01-1.0010.31670## platelets
2.502e-072.293e-06-0.1090.91313
## serum_creatinine
                                     3.950e-01 2.313e-01
                                                                 1.708 0.08767.
                                 -7.384e-024.764e-02-1.5500.12117
## serum sodium
## sex1
                                  -8.848e-015.098e-01-1.7360.08262.
                                  -3.864e-015.298e-01-0.7290.46584
## smoking1
## time
                                            -2.112e-023.921e-03-5.3877.15e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
## (Dispersion parameter for binomial family taken to be 1) ##
        Null deviance: 259.93 on 207 degrees of freedom ## Residual
deviance: 147.47 on 195 degrees of freedom
## AIC: 173.47
##
## Number of Fisher Scoring iterations: 6
predicted <- predict(mylogit_3, test_data, type="response")</pre>
```

We note a key difference here - we do not see the serum_creatinine as a significant variable in this iteration. We see only age, time and ejection_fraction as significant variables.

Deciding on optimal cutoff

```
optCutOff <- optimalCutoff(test_data$DEATH_EVENT, predicted)[1] optCutOff
```

##[1]0.6984528

This tells us that we can use this prob cutoff to classify an observation as 0 or 1. If the prob-value is below this threshold we can classify it as survival else death event.

Mis-classification error

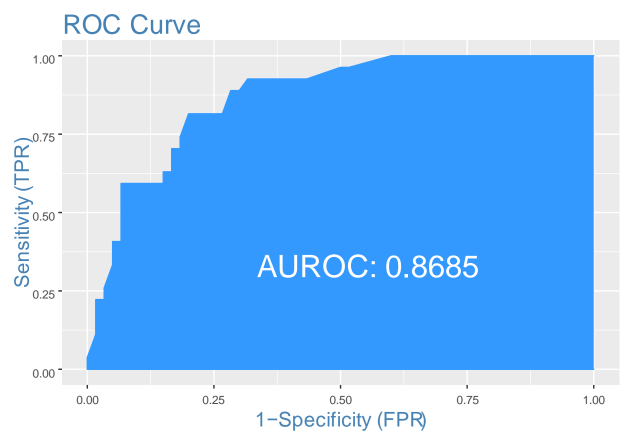
```
misClassError(test_data$DEATH_EVENT, predicted, threshold = optCutOff)
```

##[1]0.1724

We note a misclassification error on test set of 17.24%

ROC curve

plotROC(test_data\$DEATH_EVENT, predicted)

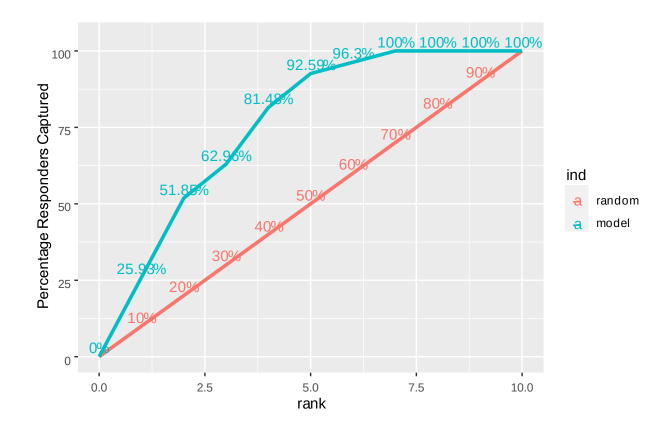


We see an AUC of 0.868 which is decent.

KS plot

```
ks_stat(test_data$DEATH_EVENT, predicted)
## [1] 0.5926
ks_plot(test_data$DEATH_EVENT, predicted)
```

KS Plot



Concordance check

Concordance(test_data\$DEATH_EVENT, predicted)

```
## $Concordance
## [1] 0.8685185
##
## $Discordance
## [1] 0.1314815
##
## $Tied
## [1] 5.551115e-17
##
## $Pairs
## [1] 1620
```

Usually concordance is in-line with AUC and we see that 86.8% pairs are concordant (the model calculated prob scores of 1s being greater than model calculated prob scores of 0s)

Using optimal cutoff to determine accuracy measures

```
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit_3,test_data,type="response")>threshold,1,0)
actual_values<-test_data$DEATH_EVENT
```

```
confusionMatrix(data = as.factor(predicted_values),
       reference = as.factor(actual_values), mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction 0
## 0 56
## 1 4##
                                                                                      Error! Bookmark not defined.
##
                       Accuracy: 0.8276
##
                           95% CI: (0.7316, 0.9002)
##
           No Information Rate: 0.6897
##
            P-Value [Acc > NIR]: 0.002661
##
                           Kappa: 0.5663
##
##
## Mcnemar's Test P-Value: 0.121335
##
##
        Precision: 0.8358##
                                 Recall:
0.9333
##
                              F1:0.8819
##
        Prevalence: 0.6897 ## Detection
Rate: 0.6437
##
        Detection Prevalence: 0.7701##
        Balanced Accuracy: 0.7630
##
##
                'Positive' Class: 0
##
We see that on test set our accuracy is 82.7%
Using optimal cutoff to determine accuracy measures with positive class as 1
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit_3, test_data, type="response")>threshold,1,0)
actual_values<-test_data<br/>
SDEATH_EVENT
confusionMatrix(data = as.factor(predicted_values), reference =
                   as.factor(actual values), positive='1', mode =
                   "prec recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction 0 1
        05611##
##
        1416
##
##
                        Accuracy: 0.8276
                           95% CI: (0.7316, 0.9002)
##
```

8

19

No Information Rate: 0.6897

##

P-Value [Acc > NIR]: 0.002661

##

Kappa: 0.5663

##

Mcnemar's Test P-Value: 0.121335

##

Precision: 0.8000## Recall:

0.5926

F1:0.6809 ## Prevalence:0.3103 ## Detection

Rate: 0.1839

Detection Prevalence : 0.2299 ## Balanced Accuracy : 0.7630

##

'Positive' Class: 1

##

We however note a key difference in this accuracy. Our recall has fallen to 0.59 while the precision is 0.80 with F1-score of 0.68

Iteration - 4 Stepwise regression

We now perform a stepwise regression which computes a null model and a full model first and adds variables as long as the added variable's AIC is below the previous computation of AIC.

```
null_model<-glm(DEATH_EVENT~1,data=train_data,family='binomial') full_model<-
glm(DEATH_EVENT~.,data=train_data,family='binomial') step_model <- step(null_model,
                       scope = list(lower = null model, upper =
                                      full model),
                       direction = "forward")
## Start: AIC=261.93
## DEATH_EVENT~1
##
##
                                    Df Deviance
                                                      AIC
## + time
                                            195.18 199.18
## + ejection fraction
                                    1
                                           235.92 239.92
## + serum creatinine
                                    1
                                            237.52 241.52
                                    1
## + age
                                           244.83 248.83
## + serum sodium
                                    1
                                           252.47 256.47
## + anaemia
                                    1
                                           257.23 261.23
## < none>
                                            259.93 261.93
## + high blood pressure
                                     1
                                           258.97 262.97
## + creatinine_phosphokinase 1
                                           259.29 263.29
## + smoking
                                            259.37 263.37
                                     1
## + sex
                                    1
                                            259.49 263.49
## + platelets
                                            259.74 263.74
                                    1
## + diabetes
                                    1
                                            259.76 263.76
##
## Step: AIC=199.18
## DEATH_EVENT ~ time
##
##
                                    Df Deviance
                                                      AIC
                                    1
                                            172.94 178.94
## + ejection_fraction
## + serum creatinine
                                    1
                                            184.74 190.74
## + serum sodium
                                    1
                                            188.05 194.05
## + age
                                    1
                                            191.07 197.07
## + smoking
                                    1
                                            192.53 198.53
## < none >
                                            195.18 199.18
## + sex
                                    1
                                            193.86 199.86
                                    1
## + high_blood_pressure
                                            194.86 200.86
                                    1
## + diabetes
                                            194.91 200.91
## + anaemia
                                    1
                                            195.00 201.00
## + platelets
                                     1
                                            195.00 201.00
## + creatinine phosphokinase 1
                                           195.10 201.10
##
## Step: AIC=178.94
## DEATH EVENT~ time + ejection fraction ##
                                    Df Deviance
                                                      AIC
## + serum_creatinine
                                            163.51 171.51
                                    1
```

```
## + age
                                   1
                                           164.56 172.56
## + serum_sodium
                                   1
                                          168.52 176.52
## + sex
                                    1
                                         168.68 176.68
                                       168.76 176.76
## + smoking
## < none >
                                         172.94 178.94
## + anaemia
                                         171.98 179.98
## + diabetes
                                    1
                                          172.38 180.38
## + high blood pressure1
                                172.50 180.50##+
creatinine_phosphokinase1
                                172.85 180.85
## + platelets
                                    1
                                          172.93 180.93
##
## Step: AIC=171.51
## DEATH_EVENT ~ time + ejection_fraction + serum_creatinine
##
##
                                    Df Deviance
                                                     AIC
## + age 1
                157.00 167.00 ## + sex 1
                                                160.28
170.28 ##+ smoking
                                161.07 171.07##+
                        1
serum sodium 1
                        161.22 171.22
## < none >
                163.51 171.51## + anaemia
        162.06 172.06
## + creatinine_phosphokinase 1
                                          163.03 173.03
## + diabetes
                                          163.05 173.05
                                    1
## + high blood pressure
                                    1
                                          163.41 173.41
## + platelets
                                          163.50 173.50
##
## Step: AIC=166.99
## DEATH_EVENT~ time + ejection_fraction + serum_creatinine + age
##
##
                                    Df Deviance
                                                     AIC
                152.78 164.78 ## + smoking
## + sex 1
                                                1
        154.68 166.68 ## + serum_sodium 1
                                                154.78
166.78
                157.00 167.00 ## + anaemia
## < none>
        155.71 167.71## + diabetes
                                        1
                                                156.09
168.09
## + creatinine_phosphokinase 1 156.79 168.79 ## +
high blood pressure
                                156.81 168.81
## + platelets
                                    1
                                          156.93 168.93
##
## Step: AIC=164.78
## DEATH EVENT~ time + ejection fraction + serum creatinine + age +
##
        sex
##
##
                                    Df Deviance
                                                     AIC
## + serum_sodium
                                          150.14 164.14
## <none>
                                          152.78 164.78
## + high blood pressure
                                          151.81 165.81
```

```
## + anaemia
                                      1
                                           152.07 166.07
## + smoking
                                      1
                                           152.08 166.08
## + diabetes
                                           152.20 166.20
                                      1
## + platelets
                                      1
                                           152.61 166.61
## + creatinine phosphokinase 1
                                           152.75 166.75
##
## Step: AIC=164.14
## DEATH_EVENT ~ time + ejection_fraction + serum_creatinine + age +
##
         sex + serum_sodium
##
                                     Df Deviance
##
                                                      AIC.
## <none>
                                           150.14 164.14
## + high blood pressure
                                           149.20 165.20
## + anaemia
                                      1
                                           149.34 165.34
## + smoking
                                      1
                                           149.53 165.53
## + diabetes
                                      1
                                           149.91 165.91
## + platelets
                                           150.05 166.05
                                      1
## + creatinine phosphokinase 1
                                           150.13 166.13
```

We see the results of the stepwise regression lowering our AIC to 164.14 with variables like time, ejection_fraction, serum_creatinine, age, sex and serum_sodium

summary(step_model)

```
##
## Call:
## glm(formula = DEATH_EVENT~ time + ejection_fraction + serum_creatinine +
               age + sex + serum_sodium, family = "binomial", data = train_data)
## Deviance Residuals:
##
       Min
               10
                        Median 3Q
                                       Max ## -1.9074 -
## Coefficients:
                              Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                        12.185514
                                      6.446110
                                                     1.890 0.05871.
                                         0.003657 -5.560 2.70e-08 * * *
                         -0.020332
## ejection fraction -0.096216
                                         0.021094 -4.561 5.08e-06 * * *
## serum creatinine
                                      0.215623
                          0.451644
                                                    2.095 0.03621*
                                      0.018369
## age
                          0.048070
                                                    2.617 0.00888 **
## sex1
                                         0.453200 -2.116 0.03435 *
                         -0.958956
                                         0.045182 -1.639 0.10120
## serum sodium
                         -0.074058
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
##
## (Dispersion parameter for binomial family taken to be 1) ##
        Null deviance: 259.93 on 207 degrees of freedom ## Residual
deviance: 150.14 on 201 degrees of freedom
## AIC: 164.14
##
## Number of Fisher Scoring iterations: 6
 We see a lower AIC but unfortunately serum_sodium isn't significant. We can remove this variable and re-compute
the ideal model.
# Model 4
mylogit_4 <- glm(DEATH_EVENT ~ time + ejection_fraction + serum_creatinine + age + sex, data
    = train_data, family = "binomial")
summary(mylogit_ 4)
##
## Call:
## glm(formula = DEATH_EVENT~ time + ejection_fraction + serum_creatinine +
##
               age + sex, family = "binomial", data = train data)
##
## Deviance Residuals:
##
                1Q
                         Median 3Q
                                         Max ## -1.9095 -
        Min
## Coefficients:
                                Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                           1.939576
                                        1.357212
                                                       1.429 0.15298
                          -0.019475
                                           0.003506 -5.554 2.79e-08 * * *
## time
## ejection_fraction -0.096797
                                           0.020669 -4.683 2.82e-06 ***
## serum creatinine
                                        0.221317
                                                       2.304 0.02120 *
                           0.509987
                                        0.018083
                                                       2.635 0.00841 **
## age
                           0.047654
## sex1
                          -0.902833
                                           0.445665 -2.026 0.04278 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
## (Dispersion parameter for binomial family taken to be 1) ##
        Null deviance: 259.93 on 207 degrees of freedom ## Residual
deviance: 152.78 on 202 degrees of freedom
## AIC: 164.78
## Number of Fisher Scoring iterations: 6
We note our lowest AIC yet of 164.78
```

Let's predict the outcome for this model

```
predicted <- predict(mylogit_4, test_data, type="response")</pre>
```

Deciding on optimal cutoff

 $optCutOff <- \ optimal Cutoff (test_data \ DEATH_EVENT, predicted) [1] \ optCutOff$

##[1]0.5069049

This tells us that we can use this prob cutoff to classify an observation as 0 or 1. If the prob-value is below this threshold we can classify it as survival else death event.

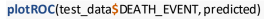
Mis-classification error

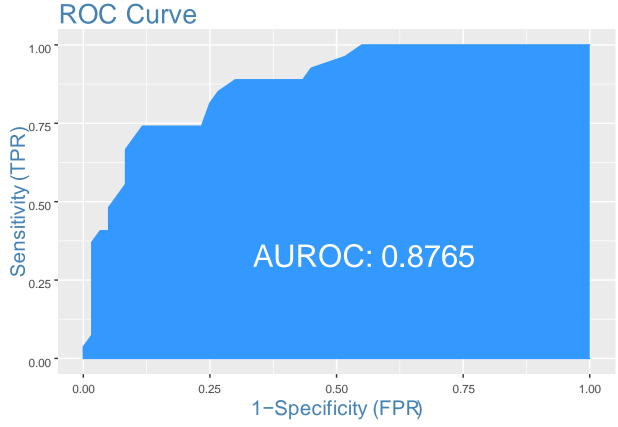
misClassError(test_data\$DEATH_EVENT, predicted, threshold = optCutOff)

##[1]0.1494

We note a misclassification error on test set of 14.9%

ROC curve



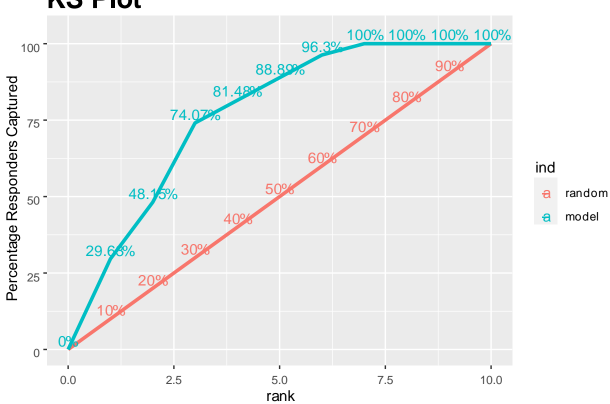


We see an AUC of 0.876 which is our best yet on test data.

KS plot

```
ks_stat(test_data$DEATH_EVENT, predicted)
##[1]0.6241
ks_plot(test_data$DEATH_EVENT, predicted)
```





Concordance check

Concordance(test_data\$DEATH_EVENT, predicted)

```
## $Concordance
##[1]0.8765432
##
## $Discordance
##[1]0.1234568
##
## $Tied ##
[1]0
##
## $Pairs
##[1]1620
```

Usually concordance is in-line with AUC and we see that 87.6% pairs are concordant (the model calculated prob scores of 1s being greater than model calculated probscores of 0s)

Using optimal cutoff to determine accuracy measures

```
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit_4,test_data, type="response")>threshold,1,0)
actual values<-test data<a href="mailto:detactual">DEATH EVENT</a>
confusionMatrix(data = as.factor(predicted values), reference =
                   as.factor(actual_values), mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction 0 1
##
                0558
##
                1519
##
                        Accuracy: 0.8506
##
                            95% CI: (0.758, 0.918)
##
            No Information Rate: 0.6897
##
##
            P-Value [Acc > NIR]: 0.0004584
##
##
                           Kappa: 0.6399
## Mcnemar's Test P-Value: 0.5790997
##
        Precision: 0.8730##
##
                                  Recall:
0.9167
##
                               F1:0.8943
##
        Prevalence: 0.6897 ## Detection
Rate: 0.6322
##
        Detection Prevalence: 0.7241##
        Balanced Accuracy: 0.8102
##
##
                'Positive' Class: 0
##
We see that on test set our accuracy is 85.0%
```

we see that off test set our accuracy is 65.0%

Using optimal cutoff to determine accuracy measures with positive class as 1

```
threshold=optCutOff

predicted_values<-ifelse(predict(mylogit_4, test_data, type="response")>threshold,1,0)

actual_values<-test_data$DEATH_EVENT confusionMatrix(data =

as.factor(predicted_values), reference = as.factor(actual_values), positive='1',mode =

"prec_recall")
```

Confusion Matrix and Statistics

##

Reference

Prediction 0 1

0558 ## 1519

##

Accuracy: 0.8506

95% CI: (0.758, 0.918)

No Information Rate : 0.6897 ## P-Value [Acc > NIR] : 0.0004584

##

Kappa: 0.6399

##

Mcnemar's Test P-Value: 0.5790997

##

Precision: 0.7917## Recall:

0.7037

F1:0.7451 ## Prevalence:0.3103 ## Detection

Rate: 0.2184

Detection Prevalence : 0.2759 ## Balanced Accuracy : 0.8102

##

'Positive' Class: 1

##

We see improved values of precision to 0.79, recall to 0.70 and F1 score to 0.74

This clearly is our most balanced and best model yet.

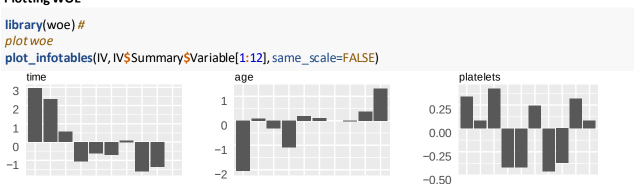
Iteration 5 - Computing WOE (weight of evidence) and IV (information value) to improve prediction accuracy

Computing IV

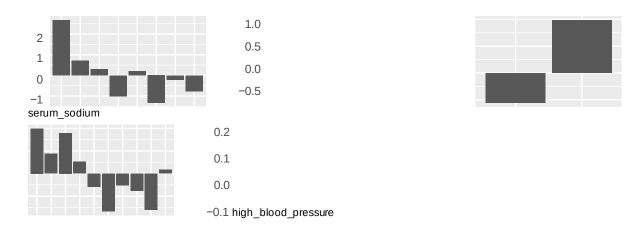
```
library(Information) library(gridExtra)
data <- read.csv' / Users/mac/Downloads/heart failure clinical records dataset.csv' ) data <-
data[datasejection fraction < 70,] data <- data[datascreatinine phosphokinase < 7000,] datasanaemia <-
factor(data$anaemia) data$diabetes<-factor(data$diabetes)
data$high blood pressure <- factor(data$high blood pressure)data$sex <-
factor(data$sex) data$smoking <- factor(data$smoking)</pre>
# this package needs the dependent variable in numeric format
# hence we reload data here
IV <- create_infotables(data=data, y="DEATH_EVENT", bins=10,
                   parallel=FALSE)
IV_Value = data.frame(IV$Summary)
IV$Summary
##
                           Variable
## 12
                                time 1.840224e+00
##5
        ejection fraction 9.763676e-01##8
        serum creatinine 9.235629e-01
                                 age 4.849249e-01
## 1
## 9
                      serum sodium 4.030774e-01
## 3 creatinine_phosphokinase 2.157046e-01
                           platelets 1.132326e-01
## 6
               high_blood_pressure 2.194425e-02
        anaemia 2.158339e-02 ##10
##2
3.028463e-04##11
                         smoking 7.862779e-05 ##4
        diabetes 8.479320e-06
```

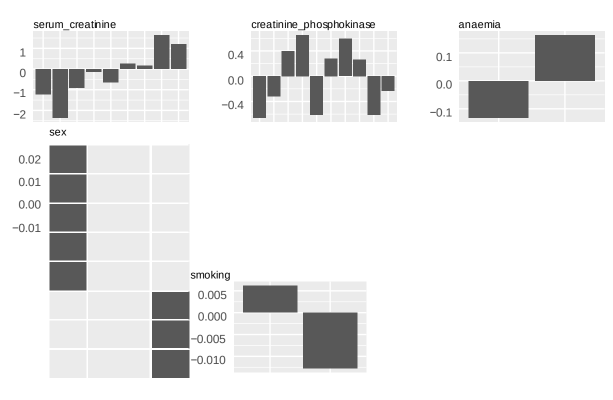
Let's analyze IV first - Our IV values are significant for time, ejection_fraction, serum_creatinine, age, serum_sodium, creatinine_phosphokinase and platelets (>0.1). After platelets, IV values are below 0.02 so we do not need to use these variables.

Plotting WOE

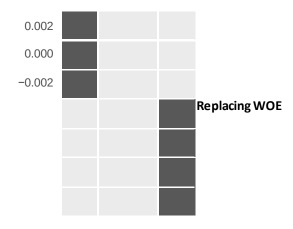


ejection_fraction





diabetes



) {

```
library(fuzzyjoin)
woe replace <- function(df orig, IV) { df <-
  cbind(df_orig)
  df_clmtyp <- data.frame(clmtyp = sapply(df, class)) df_col_typ
  <data.frame(clmnm = colnames(df), clmtyp = df clmtyp$clmtyp)
  for (rownmin 1:nrow(df col typ)) { colmn nm <-
    toString(df col typ[rownm, "clmnm"]) if(colmn nm%in%
    names(IV$Tables)){
    column_woe_df<- cbind(data.frame(IV$Tables[[toString(df_col_typ[rownm, "clmnm"])]])) if (df_col_typ[rown
    "clmtyp"] == "factor" | df col typ[rownm, "clmtyp"] == "character"
      df<-
         dplyr::inner join(df,
           column_woe_df[,c(colmn_nm,"WOE")], by
           = colmn nm, type = "inner", match = "all"
         ) df[colmn nm]<-NULL
       colnames(df)[colnames(df)=="WOE"]<-colmn nm
    } else if (df col typ[rownm, "clmtyp"] == "numeric" | df col typ[rownm, "clmtyp"] == column woe df$b<-
       as.numeric(str_sub(column woe df[,colmn nm],
         regexpr("\\[", column_woe_df[,colmn_nm]) + 1, regexpr(",",
         column_woe_df[,colmn_nm]) - 1
      ))
      column_woe_df$uv<-as.numeric(str_sub(column_woe_df[,colmn_nm],
         regexpr(",", column_woe_df[,colmn_nm]) + 1, regexpr("\\]",
         column_woe_df[,colmn_nm]) - 1
      ))
      column woe df[colmn nm]<-NULL
       column_woe_df<-column_woe_df[,c("lv","uv","WOE")]
       colnames(df)[colnames(df)==colmn nm]<-"WOE temp2381111111111111697" df <-
         fuzzy_inner_join(df,
           column_woe_df[,c("lv","uv","WOE")],
           by = c("WOE_temp238111111111111111697"="lv","WOE_temp238111111111111111697"="lv"),
           match fun=list('>=','<=')
         ) df["WOE_temp238111111111111697"]<-NULL
       df["lv"]<-NULL df["uv"]<-NULL
       colnames(df)[colnames(df)=="WOE"]<-colmn nm
    }}
  }
  return(df)
df woe <- woe replace (data, IV)
```

"integer") {

```
str(df woe)
## 'data.frame':
                            295 obs. of 13 variables:
## $ DEATH EVENT
                                           : int 111111111...
## $ age
                                          : num 0.3879 0.00248 -0.32294 0.00248 1.32221 ...
## $ anaemia
                                          : num -0.132 -0.132 0.163 0.163 0.163 ...
##$ creatinine phosphokinase: num 0.265 0.651 0.401 0.651 -0.659 ...
## $ diabetes
                                                  : num 0.00248 0.00248 0.00248 -0.00342 0.00248 ...
##$ejection fraction
                                    : num 2.67 2.67 2.67 -1.33 ...
##$high blood pressure
                                    : num 0.197 -0.112 -0.112 -0.112 0.197 ...
##$ platelets
                                    : num -0.4565 0.0825 0.4274 0.3161 0.4274 ...
##$ serum creatinine
                                    : num 1.641 0.157 1.641 1.206 1.206 ...
##$ serum sodium
                                    : num 0.999 0.999 -0.834 0.999 0.439 ...
##$ sex
                                    : num -0.0128 -0.0128 -0.0128 0.0237 -0.0128 ...
##$ smoking
                                    : num 0.00615 -0.01279 0.00615 0.00615 -0.01279 ...
                                    : num 2.94 2.94 2.94 2.94 2.94 ...
##$ time
Let's now use the new dataframe for prediction.
Splitting into train and test - 70%, 30% split
set.seed(123)
trainIndex <- createDataPartition(df woe $DEATH EVENT, p = .7,
                                           list = FALSE,
                                         times = 1
train data<-df woe[trainIndex,] test data<-df woe[_trainIndex,]
# Model 5
mylogit 5 <- glm(DEATH EVENT ~ age+anaemia+creatinine phosphokinase+
     diabetes+ejection_fraction+high_blood_pressure+platelets+
     serum creatinine+serum sodium+sex+smoking+time,
     data = train data, family = "binomial")
summary(mylogit 5)
##
## Call:
## glm(formula = DEATH_EVENT ~ age + anaemia + creatinine_phosphokinase +
                 diabetes + ejection fraction + high blood pressure + platelets +
                 serum creatinine + serum_sodium + sex + smoking + time, family = "binomial",
##
##
          data = train_data)
##
## Deviance Residuals:
                         Median 3Q
                                         Max ## -2.38680 -
                1Q
0.27533-0.052670.044662.86971
##
## Coefficients:
                                       Estimate Std. Error zvalue Pr(>|z|)
                                                     0.3805 -4.6802.87e-06 ***
## (Intercept)
                                    -1.7806
                                     1.4738
                                                  0.4871
                                                              3.026 0.002481 **
## age
```

0.7736

-1.0069

2.6262

anaemia

creatinine_phosphokinase

2.2465 -0.448 0.654019

3.395 0.000687 ***

```
## diabetes
                                    41.5959
                                                107.3797
                                                             0.387 0.698481
## ejection fraction
                                    1.8307
                                                0.4350
                                                                4.209 2.57e-05 * **
## high_blood_pressure
                                    0.1975
                                                2.1164
                                                             0.093 0.925659
## platelets
                                    0.9531
                                                0.9779
                                                             0.975 0.329764
## serum creatinine
                                    0.6848
                                                0.3016
                                                              2.271 0.023159 *
## serum sodium
                                    1.7529
                                                0.5805
                                                                 3.020 0.002529 **
## sex
                                    3.5083
                                                18.9358
                                                             0.185 0.853012
## smoking
                                  -45.4997
                                                38.7693 - 1.174 0.240555
## time
                                    1.6880
                                                0.3265
                                                                5.171 2.33e-07 * **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
## (Dispersion parameter for binomial family taken to be 1) ##
        Null deviance: 251.006 on 206 degrees of freedom ## Residual
deviance: 80.756 on 194 degrees of freedom
## AIC: 106.76
##
## Number of Fisher Scoring iterations: 7
predicted <- predict(mylogit_5, test_data, type="response")</pre>
We see age, time, ejection_fraction and serum_sodium, creatinine_phosphpkinase, serum_creatinine as significant
variables.
# Model 5
mylogit_5 <- glm(DEATH_EVENT ~ age + ejection_fraction+
     serum sodium+time+creatinine phosphokinase+ serum creatinine, data =
         train data, family = "binomial")
summary(mylogit 5)
##
## Call:
## glm(formula = DEATH_EVENT ~ age + ejection_fraction + serum_sodium +
##
                 time + creatinine_phosphokinase + serum_creatinine, family = "binomial",
##
          data = train_data)
##
## Deviance Residuals:
                                          Max ## -2.60054 -
##
        Min
                1Q
                         Median 3Q
0.28769-0.067550.071442.75776
##
## Coefficients:
##
                                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                                        0.3457 -4.8431.28e-06 ***
                                  -1.6741
## age
                                     1.4124
                                                  0.4665
                                                                  3.028 0.00246 * *
                                     1.6660
                                                  0.3847
                                                                4.331 1.49e-05 ***
## ejection_fraction
## serum sodium
                                     1.6252
                                                  0.5245
                                                                  3.099 0.00194 * *
## time
                                                  0.2864
                                                                5.442 5.27e-08 ***
                                     1.5588
                                                                3.390 0.00070 ***
## creatinine phosphokinase
                                     2.4491
                                                  0.7225
## serum creatinine
                                     0.6359
                                                  0.2825
                                                               2.251 0.02437 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
```

```
##
## (Dispersion parameter for binomial family taken to be 1) ##
## Null deviance: 251.006 on 206 degrees of freedom ## Residual
deviance: 83.779 on 200 degrees of freedom
## AIC: 97.779
##
## Number of Fisher Scoring iterations: 7

predicted <- predict(mylogit_5, test_data, type="response")
We note our lowest AIC yet of 97.77
```

Deciding on optimal cutoff

```
optCutOff <- optimalCutoff(test_data$DEATH_EVENT, predicted)[1] optCutOff
```

##[1]0.09998813

This tells us that we can use this prob cutoff to classify an observation as 0 or 1. If the prob-value is below this threshold we can classify it as survival else death event.

Mis-classification error

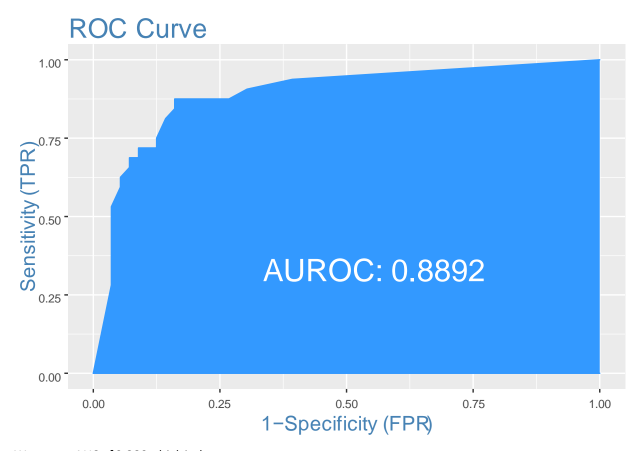
```
misClassError(test_data$DEATH_EVENT, predicted, threshold = optCutOff)
```

##[1]0.1477

We note a misclassification error on test set of 14.7%

ROC curve

plotROC(test_data\$DEATH_EVENT, predicted)

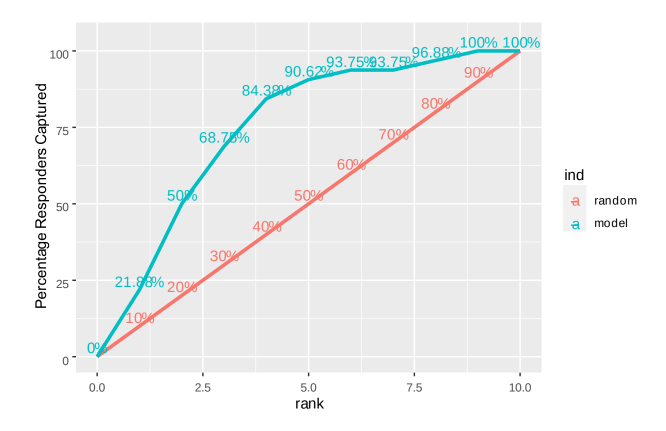


We see an AUC of 0.889 which is decent.

KS plot

```
ks_stat(test_data$DEATH_EVENT, predicted)
## [1] 0.683
ks_plot(test_data$DEATH_EVENT, predicted)
```

KS Plot



Concordance check

Concordance(test_data\$DEATH_EVENT, predicted)

```
## $Concordance
## [1] 0.8889509
##
## $Discordance
## [1] 0.1110491
##
## $Tied
## [1] -4.163336e-17
##
## $Pairs
## [1] 1792
```

Usually concordance is in-line with AUC and we see that 88.8% pairs are concordant (the model calculated prob scores of 1s being greater than model calculated prob scores of 0s)

Using optimal cutoff to determine accuracy measures

```
threshold=optCutOff
predicted_values<-ifelse(predict(mylogit_5,test_data, type="response")>threshold,1,0)
actual_values<-test_data_DEATH_EVENT
```

```
confusionMatrix(data = as.factor(predicted values), reference =
    as.factor(actual_values), mode = "prec_recall")
## Confusion Matrix and Statistics
##
##
                Reference
## Prediction 0 1
##
               0474
##
               1928
##
##
                       Accuracy: 0.8523
##
                           95% CI: (0.7606, 0.9189)
##
           No Information Rate: 0.6364
            P-Value [Acc > NIR]: 6.225e-06
##
##
##
                          Kappa: 0.6911
##
## Mcnemar's Test P-Value: 0.2673##
##
        Precision: 0.9216##
0.8393
##
                              F1:0.8785
##
        Prevalence: 0.6364 ##
                                Detection
Rate: 0.5341
##
        Detection Prevalence: 0.5795##
        Balanced Accuracy: 0.8571
##
##
               'Positive' Class: 0
##
We see that on test set our accuracy is 85.2%
```

Using optimal cutoff to determine accuracy measures with positive class as 1

Accuracy: 0.8523

No Information Rate: 0.6364

95% CI: (0.7606, 0.9189)

##

##

##

```
threshold=optCutOff

predicted_values<-ifelse(predict(mylogit_5, test_data, type="response")>threshold,1,0)

actual_values<-test_data$DEATH_EVENT confusionMatrix(data =

as.factor(predicted_values), reference = as.factor(actual_values), positive='1',mode =

"prec_recall")

## Confusion Matrix and Statistics

##

## Reference

## Prediction 0 1

## 0 47 4

## 1 9 28

##
```

P-Value [Acc > NIR]: 6.225e-06

##

Kappa: 0.6911

##

Mcnemar's Test P-Value: 0.2673##
Precision: 0.7568## Recall:

0.8750

F1:0.8116 ## Prevalence:0.3636## Detection

Rate: 0.3182

Detection Prevalence : 0.4205 ## Balanced Accuracy : 0.8571

##

'Positive' Class: 1

##

We however note a key difference in this accuracy. Our recall is 0.87 while the precision is 0.75 with F1-score of 0.81

We see how computing and recoding variables to WOE improved our model accuracy even further. We also see our highest recall yet of 0.87 which is great for the purpose of predicting death events more rigorously.

Summarizing all model results in a table

Model	Variables	AIC	AUC	Accuracy	Precision	Recall	F1-Score	Comments
Model 1	All 241.95 0.8		5 0.894	0.854	0.79	0.72	0.75	No test set (overfitting
Model 2	Age, ejec- tion_fraction, serum_creation			0.85	0.84	0.64	0.73	No test set (overfitting
Model 3	All	173.	4 0.868	0.827	0.80	0.59	0.68	Test set resultsFirst real model
Model 4	Age, ejec- tion_fraction, serum_creation time, sex		4.7 0.87	0.85	0.79	0.70	0.74	Stepwise Forward selection
Model 5	Age, ejec- tion_fraction, serum_creation serum_sodium creatinine_ph	nine, tii m,	me,	0.852	0.75	0.88	0.81	WoE dataset

)

We note that in model 4 stepwise method performed well and gave us better model results than model 3

However, we finally have a model (Model 5) which we can use as a best model outcome from our iterations which came from computing woe and iv values for each of our variables in the dataset.

This concludes our approach to Logistic regression in our dataset