Heart Failure Prediction

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Load Packages and Data

```
library(pROC)
library(caret)
library(ggplot2)
library(dplyr)
library(tidyr)
library(corrplot)
library(pRROC)
library(pRROC)
library(ggbiplot)
library(factoextra)
library(rpart.plot)

# Load data
hfp_data <- read.csv("data/heart_failure_clinical_records_dataset.csv")</pre>
```

1. Explore Dataset

```
# read structure/summary
str(hfp_data)
```

```
## 'data.frame': 299 obs. of 13 variables:
                         : num 75 55 65 50 65 90 75 60 65 80 ...
## $ age
## $ anaemia
                          : int 0001111101...
## $ creatinine phosphokinase: int 582 7861 146 111 160 47 246 315 157 123 ...
## $ diabetes
                         : int 0000100100...
  $ 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 ...
                        : num 1.9 1.1 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.4 ...
## $ serum_creatinine
## $ serum sodium
                        : int 130 136 129 137 116 132 137 131 138 133 ...
## $ sex
                        : int 1111011101...
## $ smoking
                        : int 0010010101...
                        : int 46778810101010...
## $ time
  $ DEATH EVENT
                         : int 111111111...
```

summary(hfp data)

```
##
                    anaemia
                                creatinine_phosphokinase
                                                         diabetes
        age
##
   Min. :40.00
                 Min.
                       :0.0000
                                Min.
                                      : 23.0
                                                      Min.
                                                            :0.0000
  1st Qu.:51.00
                1st Qu.:0.0000
                                1st Qu.: 116.5
                                                      1st Qu.:0.0000
##
   Median :60.00 Median :0.0000
                                Median : 250.0
                                                      Median :0.0000
   Mean :60.83 Mean :0.4314
                                Mean : 581.8
                                                     Mean :0.4181
##
   3rd Qu.:70.00
                 3rd Qu.:1.0000
                                3rd Qu.: 582.0
                                                     3rd Qu.:1.0000
##
##
   Max. :95.00
                 Max. :1.0000
                                Max. :7861.0
                                                      Max. :1.0000
   ejection_fraction high_blood_pressure platelets
                                                    serum creatinine
##
   Min.
        :14.00
                   Min. :0.0000
                                   Min. : 25100
                                                   Min. :0.500
##
##
  1st Qu.:30.00
                   1st Qu.:0.0000
                                    1st Qu.:212500
                                                   1st Qu.:0.900
   Median :38.00
                  Median :0.0000
                                 Median :262000
##
                                                   Median :1.100
##
   Mean :38.08
                   Mean :0.3512
                                   Mean :263358
                                                   Mean :1.394
   3rd Qu.:45.00
                                    3rd Qu.:303500
##
                 3rd Qu.:1.0000
                                                  3rd Qu.:1.400
                                  Max. :850000 Max. :9.400
## Max.
        :80.00
                   Max. :1.0000
##
   serum sodium
                     sex
                                  smoking
                                                   time
   Min. :113.0 Min. :0.0000 Min. :0.0000 Min. : 4.0
##
##
  1st Qu.:134.0 1st Qu.:0.0000
                                1st Qu.:0.0000 1st Qu.: 73.0
   Median :137.0 Median :1.0000
                                Median :0.0000 Median :115.0
##
   Mean :136.6 Mean :0.6488
                                Mean :0.3211 Mean :130.3
##
  3rd Qu.:140.0
##
                 3rd Qu.:1.0000
                                3rd Qu.:1.0000 3rd Qu.:203.0
        :148.0
                 Max. :1.0000
                                Max. :1.0000 Max. :285.0
##
   Max.
   DEATH EVENT
##
## Min. :0.0000
##
  1st Qu.:0.0000
## Median :0.0000
## Mean :0.3211
## 3rd Qu.:1.0000
## Max. :1.0000
```

head(hfp_data)

a <dbl></dbl>	anaemia <int></int>	creatinine_phosphokinase <int></int>	diabetes <int></int>	ejection_fraction <int></int>			
1 75	0	582	0	20			
2 55	0	7861	0	38			
3 65	0	146	0	20			
4 50	1	111	0	20			
5 65	1	160	1	20			
6 90	1	47	0	40			
6 rows I 1-6 of 14 columns							

```
# check missing values
colSums(is.na(hfp_data))
```

```
##
                                                anaemia creatinine_phosphokinase
                         age
                           0
##
                                     ejection_fraction
##
                    diabetes
                                                             high_blood_pressure
##
##
                   platelets
                                      serum_creatinine
                                                                     serum_sodium
##
                           0
                                                                                 0
                                                smoking
                                                                             time
##
                         sex
##
                                                                                 0
##
                 DEATH_EVENT
                           0
##
```

```
# distribution
table(hfp_data$DEATH_EVENT)
```

```
##
## 0 1
## 203 96
```

```
prop.table(table(hfp_data$DEATH_EVENT))
```

```
##
## 0.6789298 0.3210702
```

203 (67.8%) of patients survived, while 96 (32.1%) died.

Which variables are related to whether a patient died from heart failure (response variable (DEATH_EVENT)?

2. Exploratory Data Analysis

2. 1 Correlation Matrix

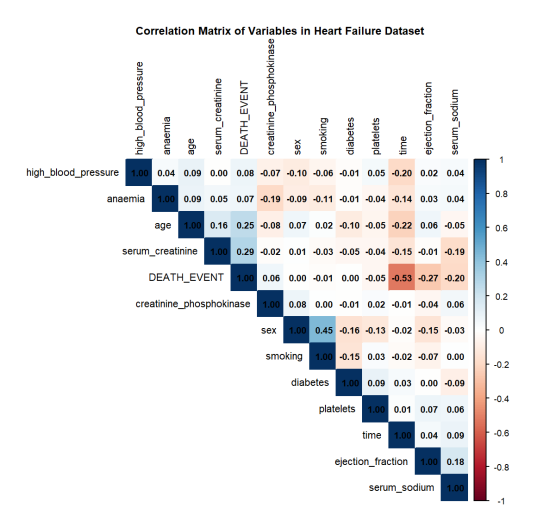
```
# correlation matrix
cor_matrix_y <- cor(hfp_data)
cor_matrix_y</pre>
```

```
##
                                         anaemia creatinine_phosphokinase
                                  age
## age
                           1.00000000 0.08800644
                                                            -0.081583900
## anaemia
                           0.08800644 1.00000000
                                                            -0.190741030
## creatinine_phosphokinase -0.08158390 -0.19074103
                                                             1.000000000
## diabetes
                          -0.10101239 -0.01272905
                                                            -0.009638514
## ejection_fraction
                           0.06009836 0.03155697
                                                            -0.044079554
## high_blood_pressure
                           0.09328868 0.03818200
                                                            -0.070589980
## platelets
                          -0.05235437 -0.04378555
                                                             0.024463389
## serum_creatinine
                           0.15918713 0.05217360
                                                            -0.016408480
## serum sodium
                          -0.04596584 0.04188161
                                                             0.059550156
## sex
                           0.06542952 -0.09476896
                                                             0.079790629
## smoking
                           0.01866787 -0.10728984
                                                             0.002421235
## time
                          -0.22406842 -0.14141398
                                                            -0.009345653
## DEATH EVENT
                           0.25372854 0.06627010
                                                             0.062728160
##
                              diabetes ejection_fraction high_blood_pressure
                                             0.06009836
## age
                          -0.101012385
                                                               0.093288685
## anaemia
                          -0.012729046
                                             0.03155697
                                                               0.038182003
## creatinine_phosphokinase -0.009638514
                                            -0.04407955
                                                              -0.070589980
## diabetes
                           1.000000000
                                            -0.00485031
                                                              -0.012732382
## ejection_fraction
                          -0.004850310
                                             1.00000000
                                                               0.024444731
## high blood pressure
                          -0.012732382
                                             0.02444473
                                                               1.000000000
## platelets
                           0.092192828
                                             0.07217747
                                                               0.049963481
## serum_creatinine
                          -0.046975315
                                            -0.01130247
                                                              -0.004934525
## serum_sodium
                          -0.089550619
                                             0.17590228
                                                               0.037109470
## sex
                          -0.157729504
                                            -0.14838597
                                                              -0.104614629
## smoking
                          -0.147173413
                                            -0.06731457
                                                              -0.055711369
## time
                           0.033725509
                                             0.04172924
                                                              -0.196439479
## DEATH EVENT
                          -0.001942883
                                            -0.26860331
                                                               0.079351058
##
                            platelets serum_creatinine serum_sodium
## age
                          -0.05235437
                                          0.159187133 -0.045965841 0.065429524
                                          0.052173604 0.041881610 -0.094768961
## anaemia
                          -0.04378555
## creatinine phosphokinase 0.02446339
                                        -0.016408480 0.059550156 0.079790629
                           0.09219283 -0.046975315 -0.089550619 -0.157729504
0.07217747 -0.011302475 0.175902282 -0.148385965
## diabetes
## ejection_fraction
## high_blood_pressure
                                        0.04996348
## platelets
                                       -0.041198077 0.062124619 -0.125120483
                           1.00000000
                                          1.000000000 -0.189095210 0.006969778
## serum_creatinine
                          -0.04119808
## serum_sodium
                           0.06212462
                                        -0.189095210 1.000000000 -0.027566123
## sex
                          -0.12512048
                                          0.006969778 -0.027566123 1.000000000
                           0.02823445 -0.027414135 0.004813195 0.445891712
## smoking
## time
                           0.01051391
                                        ## DEATH_EVENT
                          -0.04913887
                                          0.294277561 -0.195203596 -0.004316376
##
                                              time DEATH_EVENT
                               smoking
                           0.018667868 -0.224068420 0.253728543
## age
## anaemia
                          -0.107289838 -0.141413982 0.066270098
## creatinine_phosphokinase 0.002421235 -0.009345653 0.062728160
## diabetes
                          ## ejection_fraction
                          ## high_blood_pressure
                          -0.055711369 -0.196439479 0.079351058
## platelets
                           ## serum creatinine
                          -0.027414135 -0.149315418 0.294277561
                           0.004813195 0.087640000 -0.195203596
## serum_sodium
```

```
# top predictors most correlated with DEATH_EVENT
cor_target <- cor_matrix_y["DEATH_EVENT", ]
top_5 <- sort(abs(cor_target[names(cor_target) != "DEATH_EVENT"]), decreasing = TRUE)[1:5]
top_5</pre>
```

```
## time serum_creatinine ejection_fraction age
## 0.5269638 0.2942776 0.2686033 0.2537285
## serum_sodium
## 0.1952036
```

```
# heatmap
corrplot(cor_matrix_y,
        method = "color",
        type = "upper",
                             # upper triangle
        order = "hclust",
                              # group similar variables
        addCoef.col = "black", # show correlation values
        tl.cex = 0.7,
                              # axis text
        number.cex = 0.6,
                             # correlation values
        tl.col = "black",
                             # axis label color
        cl.cex = 0.6,
                              # color legend text size
        mar = c(1, 1, 2.5, 1)) # plot margins
title("Correlation Matrix of Variables in Heart Failure Dataset", cex.main = 0.75)
```



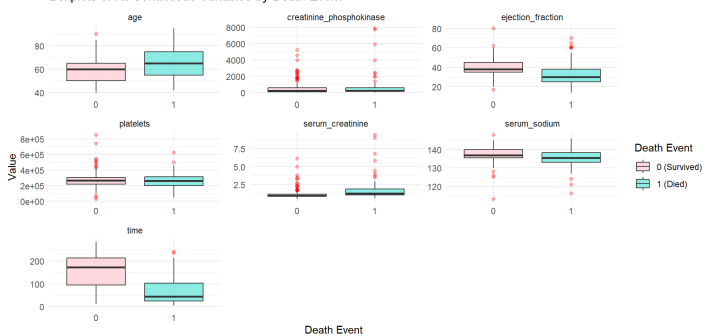
From this matrix, we can see that the top 5 predictors most correlated with <code>DEATH_EVENT</code> are time, <code>serum_creatinine</code>, <code>ejection_fraction</code>, <code>age</code>, and <code>serum_sodium</code>.

2.2 Boxplots of Continuous Features

To observe visual differences, we explored all continuous variables grouped by <code>DEATH_EVENT</code> .

```
# ID binary vars and derive continuous ones
binary_vars <- c("sex", "diabetes", "high_blood_pressure", "smoking", "anaemia")</pre>
continuous_vars <- setdiff(names(hfp_data), c(binary_vars, "DEATH_EVENT"))</pre>
# include all continuous vars + DEATH_EVENT
box_vars <- hfp_data[, c(continuous_vars, "DEATH_EVENT")]</pre>
# pivot longer for faceted plotting
long_box <- pivot_longer(box_vars,</pre>
                          cols = -DEATH_EVENT,
                          names_to = "Variable",
                          values_to = "Value")
# plot
ggplot(long_box, aes(x = factor(DEATH_EVENT), y = Value, fill = factor(DEATH_EVENT))) +
  geom_boxplot(alpha = 0.6, outlier.color = "red", outlier.alpha = 0.3) +
  facet wrap(~ Variable, scales = "free", ncol = 3) +
  labs(title = "Boxplots of All Continuous Variables by Death Event",
       x = "Death Event", y = "Value", fill = "Death Event") +
  scale_fill_manual(values = c("0" = "pink", "1" = "turquoise"),
                    labels = c("0 (Survived)", "1 (Died)")) +
  theme_minimal()
```

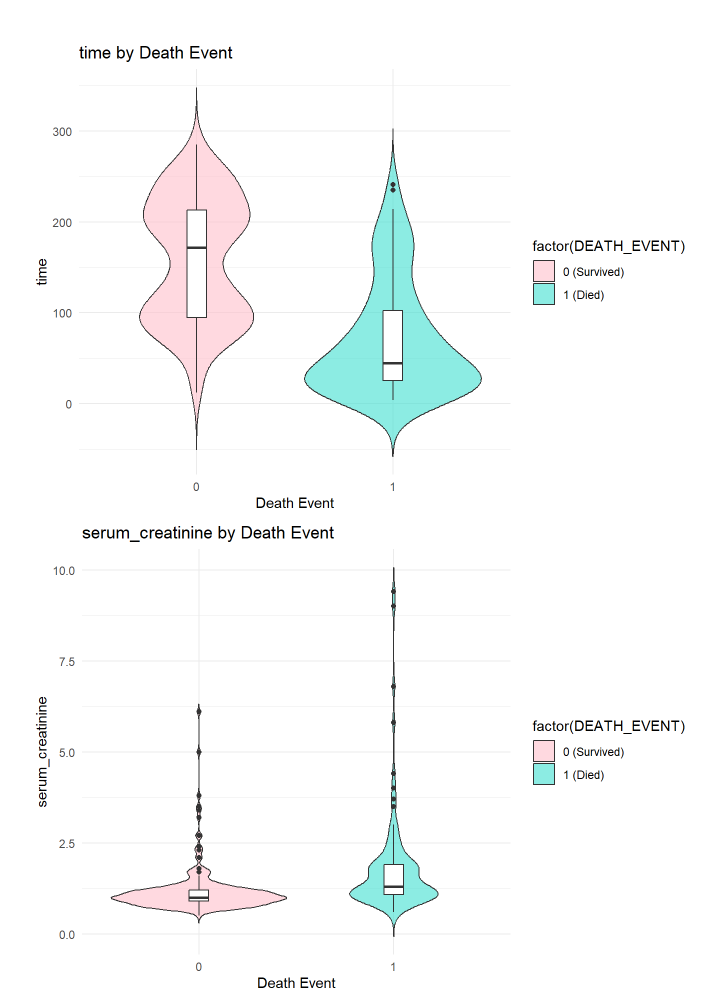
Boxplots of All Continuous Variables by Death Event



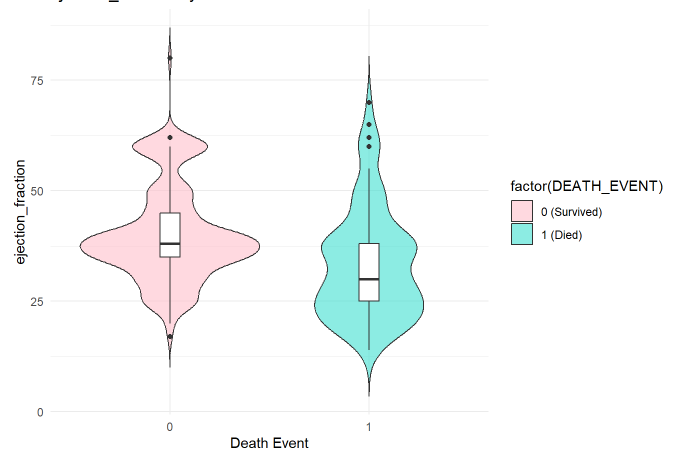
These boxplots align with the top predictors seen in the correlation matrix above.

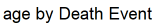
2.3 Violin Plots for Top Predictors

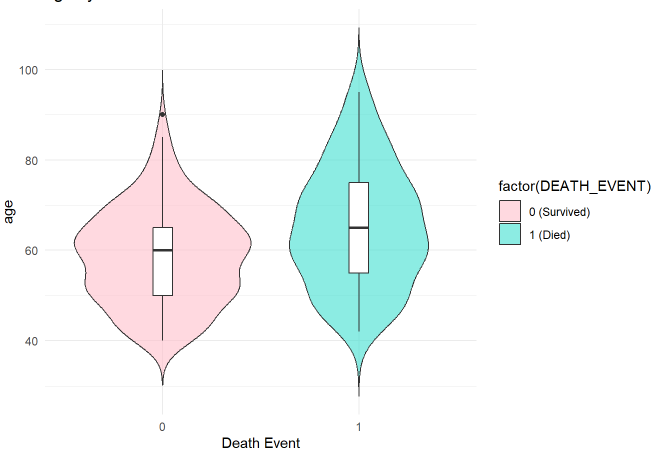
Based on the correlation and boxplots, we've selected the top 5 features to explore more deeply.



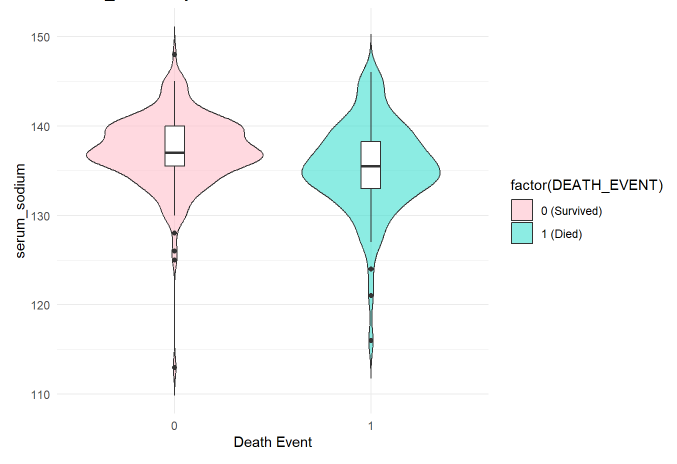
ejection_fraction by Death Event







serum sodium by Death Event

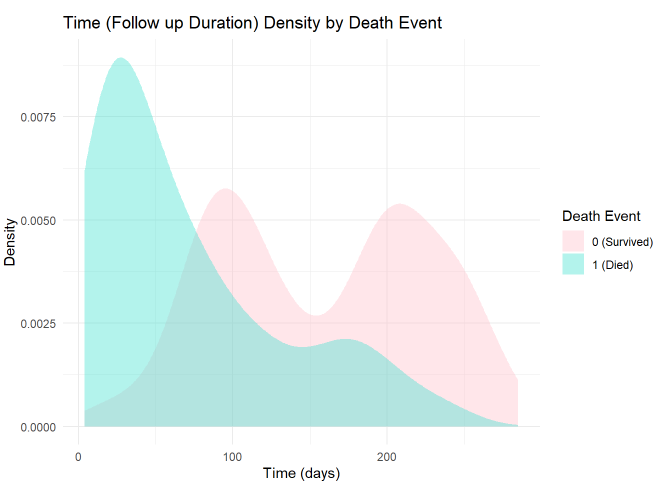


Each of these plots shows the value comparision of partients who survived (DEATH_EVENT =0), and died (DEATH_EVENT =1)

- time: patients who survived had a longer time where they followed up compared to those who died.
- serum_creatnine: shows more right-skewed distributions for patients who died, which shows that they had higher levels of serum creatnine, an indicator of poor kidney function.
- ejection_fraction : there was more instances of lower ejection fraction (how much blood the heart pumps out each heartbeat) in patients who died.
- age: the died group skews older, confirming that age is a risk factor, but still there is some overlap so it shouldn't be used as a sole predictor.
- serum_sodium: the distributions are a bit similar, with slightly lower sodium levels in patients who died. Since they are so similar, this variable might be weakly predictive.

2.4 Density Plot for Age

We wanted to explore how the top correlated variable (time) is distributed across the response group.



This plot shows the estimated probability density of follow up-time (in days). The blue curve (deaths) is heavily concentrated at lower times (<50 days), and suggests that patients who died had shorter follow up durations. The pink curve shows a "bimodal" distribution (two peaks at ~75 and 200), and indicates a wider range and longer time duration among surviving patients. This feature may contribute more significantly to class seperation and therefore reduce error training rate in models like LDA or Logistic Regression.

3. Preprocessing

3.1 Scaling Predictors

```
binary_vars <- c("sex", "diabetes", "high_blood_pressure", "smoking", "anaemia")
continuous_vars <- setdiff(names(hfp_data), c(binary_vars, "DEATH_EVENT"))

scaled_cont <- scale(hfp_data[, continuous_vars])
binary_data <- hfp_data[, binary_vars]

hfp_scaled <- cbind(as.data.frame(scaled_cont), binary_data, DEATH_EVENT = hfp_data$DEATH_EVENT)

head(hfp_scaled)</pre>
```

	age <dbl></dbl>	creatinine_phosphokinase <dbl></dbl>	ejection_fraction <dbl></dbl>	platelets <dbl></dbl>			
1	1.1909487	0.000165451	-1.527997920	1.678834e-02			
2	-0.4904571	7.502062717	-0.007064906	7.523048e-09			
3	0.3502458	-0.449185725	-1.527997920	-1.036336e+00			
4	-0.9108085	-0.485257493	-1.527997920	-5.455595e-01			
5	0.3502458	-0.434757017	-1.527997920	6.507077e-01			
6	2.4520030	-0.551217299	0.161927651	-6.069065e-01			
6 rows I 1-5 of 14 columns							

All continuous features were standardized using z-score scaling (mean = 0, sd = 1). This makes sure that the features have equal weight in distance-based methods. The binary features were kept the same to keep their categorical interpretation.

3.2 Train-Test Split

```
hfp_data$DEATH_EVENT <- factor(hfp_data$DEATH_EVENT, levels = c(0, 1), labels = c("Survived", "D
ied"))
set.seed(2025)
train_index <- createDataPartition(hfp_scaled$DEATH_EVENT, p = 0.7, list = FALSE)
train_data <- hfp_scaled[train_index, ]
test_data <- hfp_scaled[-train_index, ]</pre>
```

The dataset was split 70/30 using stratified sampling, which makes sure that the data distribution is consistent, keeping the same amount of instances where patients have died for the training and test set.

3.3 K-Fold Cross Validation

```
train_control <- trainControl(method="cv", number=10, classProbs = TRUE, summaryFunction = twoCl
assSummary, savePredictions = TRUE)</pre>
```

10-fold cross validation was set up for all models.

4. Model Training

We trained the following classifiers:

- Quadratic Discriminant Analysis (QDA)
- Linear Discriminant Analysis (LDA)
- Logistic Regression (LogReg)
- k-Nearest Neighbors (KNN)
- Support Vector Machine (SVM)
- Random Forest (RF)
- Decision Tree (DT)

Since there are binary values, we performed a conversion in the beginning to designate values for patients who died and survived.

```
#Classfication Models
# Convert DEATH_EVENT to a factor for classification since it is numeric (meant for regression)
#test_data$DEATH_EVENT <- as.factor(test_data$DEATH_EVENT)
#train_data$DEATH_EVENT <- as.factor(train_data$DEATH_EVENT)

test_data$DEATH_EVENT <- factor(test_data$DEATH_EVENT, levels = c(0, 1), labels = c("Survived",
"Died"))
train_data$DEATH_EVENT <- factor(train_data$DEATH_EVENT, levels = c(0, 1), labels = c("Survived",
"Died"))

# QDA Model
set.seed(2025)
qda_model <- train(DEATH_EVENT ~ ., data = train_data, method = "qda", trControl = train_control, metric = "ROC")
print(qda_model)</pre>
```

```
## Quadratic Discriminant Analysis
##
## 210 samples
   12 predictor
     2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 188, 190, 189, 189, ...
## Resampling results:
##
     ROC
##
                Sens
                           Spec
##
     0.7677778 0.8928571 0.447619
```

```
# LDA Model
lda_model <- train(DEATH_EVENT ~ ., data = train_data, method = "lda", trControl = train_contro
l, metric = "ROC")
print(lda_model)</pre>
```

```
## Linear Discriminant Analysis
##
## 210 samples
## 12 predictor
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 188, 189, 190, 189, 189, 190, ...
## Resampling results:
##
    ROC
##
                Sens
                           Spec
##
    0.8396032 0.8980952 0.6404762
```

```
# Logistic Regression
logistic_regression_model <- train(DEATH_EVENT ~ ., data = train_data, method = "glm", family =
"binomial", trControl = train_control, metric = "ROC")
print (logistic_regression_model)</pre>
```

```
## Generalized Linear Model
##
## 210 samples
## 12 predictor
##
    2 classes: 'Survived', 'Died'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 190, 188, 189, 189, ...
## Resampling results:
##
##
    ROC
                Sens
                           Spec
##
    0.8349206 0.8909524 0.6142857
```

```
# KNN
knn_model <- train(DEATH_EVENT ~ ., data = train_data, method = "knn", trControl = train_contro
l, metric = "ROC")
print (knn_model)</pre>
```

```
## k-Nearest Neighbors
##
## 210 samples
## 12 predictor
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 189, 188, 188, 189, ...
## Resampling results across tuning parameters:
##
##
    k ROC
                   Sens
                             Spec
##
   5 0.7699206 0.9533333 0.3547619
   7 0.8013889 0.9466667 0.3380952
    9 0.7920238 0.9533333 0.3880952
##
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was k = 7.
```

```
# SVM
svm_model <- train(DEATH_EVENT ~ ., data = train_data, method = "svmRadial", trControl = train_c
ontrol, preProcess = c("center", "scale"), metric = "ROC")
print(svm_model)</pre>
```

```
## Support Vector Machines with Radial Basis Function Kernel
##
## 210 samples
##
  12 predictor
   2 classes: 'Survived', 'Died'
##
##
## Pre-processing: centered (12), scaled (12)
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 190, 189, 189, 190, ...
## Resampling results across tuning parameters:
##
##
    C
           ROC
                      Sens
                                 Spec
##
   0.25 0.8511111 0.8709524 0.6095238
##
    0.50 0.8535714 0.8842857 0.5928571
    1.00 0.8499206 0.8771429 0.5595238
##
##
## Tuning parameter 'sigma' was held constant at a value of 0.05627339
## ROC was used to select the optimal model using the largest value.
## The final values used for the model were sigma = 0.05627339 and C = 0.5.
```

```
# Random Forest
rf_model <- train(DEATH_EVENT ~ ., data = train_data, method = "rf", trControl = train_control,
metric = "ROC")
print(rf_model)</pre>
```

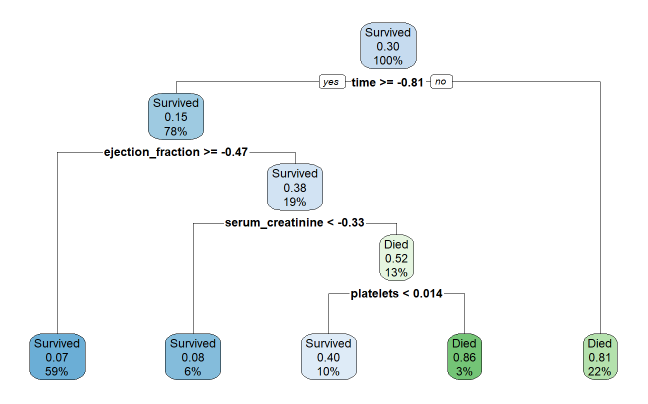
```
## Random Forest
##
## 210 samples
   12 predictor
##
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 189, 189, 189, 188, ...
## Resampling results across tuning parameters:
##
##
    mtry ROC
                      Sens
                                 Spec
##
    2
           0.8975907 0.9461905 0.5761905
##
     7
           0.8858050 0.8923810 0.6119048
##
     12
           0.8653288 0.8790476 0.5952381
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
```

```
# Decision Tree
dt_model <- train(DEATH_EVENT ~ ., data = train_data, method = "rpart", trControl = train_contro
l, metric = "ROC")
print(dt_model)</pre>
```

```
## CART
##
## 210 samples
##
   12 predictor
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 190, 188, 189, 189, 189, 188, ...
## Resampling results across tuning parameters:
##
##
                 ROC
                            Sens
                                       Spec
    ср
##
    0.01612903 0.7930952 0.8723810 0.6404762
    0.02688172 0.7898413 0.8857143 0.6547619
##
    0.46774194 0.5919048 0.9457143 0.2380952
##
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was cp = 0.01612903.
```

```
rpart.plot(dt_model$finalModel, main = "Decision Tree for Heart Failure Prediction")
```

Decision Tree for Heart Failure Prediction



dt_pred <- predict(dt_model, newdata = test_data)
confusionMatrix(dt_pred, test_data\$DEATH_EVENT)</pre>

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
                    51
##
     Survived
                         25
##
     Died
                     4
##
##
                  Accuracy : 0.8539
##
                    95% CI: (0.7632, 0.9199)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 9.176e-07
##
##
##
                     Kappa: 0.6817
##
    Mcnemar's Test P-Value: 0.2673
##
##
               Sensitivity: 0.9273
##
               Specificity: 0.7353
##
##
            Pos Pred Value: 0.8500
##
            Neg Pred Value : 0.8621
                Prevalence: 0.6180
##
            Detection Rate: 0.5730
##
      Detection Prevalence: 0.6742
##
##
         Balanced Accuracy: 0.8313
##
##
          'Positive' Class : Survived
##
```

5. Model Evaluation: F1 and AUC

5.1 Utility Functions

```
# Function to calculate F1 from confusion matrix
calculate_f1 <- function(cm) {</pre>
  precision <- cm$byClass["Pos Pred Value"]</pre>
  recall <- cm$byClass["Sensitivity"]</pre>
  # Handle division by zero
  if ((precision + recall) == 0) {
    return(NA)
  f1 <- 2 * (precision * recall) / (precision + recall)</pre>
  return(round(f1, 4))
}
calculate_auc <- function(actual, predicted_probs, positive_label = "Died") {</pre>
  actual <- factor(actual, levels = c("Survived", "Died"))</pre>
  predicted_probs <- as.numeric(predicted_probs)</pre>
  roc_curve <- roc(response = actual, predictor = predicted_probs, levels = c("Survived", "Die</pre>
d"))
  auc_value <- auc(roc_curve)</pre>
  return(round(auc_value, 4))
}
```

5.2 Metrics per Model

We generated confusion matrices for each of the models, along with manually calculating F1-score and AUC.

```
# Logistic Regression
logreg_pred <- predict(logistic_regression_model, newdata = test_data)
logreg_cm <- confusionMatrix(logreg_pred, test_data$DEATH_EVENT, positive = "Died")
logreg_f1 <- calculate_f1(logreg_cm)
print(logreg_cm)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    48
##
     Died
                     7
                         25
##
##
                  Accuracy : 0.8202
##
                    95% CI: (0.7245, 0.8936)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 3.019e-05
##
##
##
                     Kappa: 0.6149
##
   Mcnemar's Test P-Value: 0.8026
##
##
               Sensitivity: 0.7353
##
               Specificity: 0.8727
##
            Pos Pred Value : 0.7812
##
##
            Neg Pred Value : 0.8421
                Prevalence: 0.3820
##
            Detection Rate: 0.2809
##
      Detection Prevalence: 0.3596
##
##
         Balanced Accuracy: 0.8040
##
##
          'Positive' Class : Died
##
cat("\nF1: ", logreg_f1, "\n") #Calculating F1
##
## F1: 0.7576
```

```
logreg_probs <- predict(logistic_regression_model, newdata = test_data, type = "prob")</pre>
auc_value <- calculate_auc(test_data$DEATH_EVENT, logreg_probs$Died) #Calculating AUC</pre>
cat("AUC:", auc_value, "\n\n\n")
```

```
## AUC: 0.9032
```

```
# LDA
lda_pred <- predict(lda_model, newdata = test_data)</pre>
lda_cm <- confusionMatrix(lda_pred, test_data$DEATH_EVENT, positive = "Died")</pre>
lda_f1 <- calculate_f1(lda_cm)</pre>
print(lda_cm)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    48
##
     Died
                     7
                         26
##
##
                  Accuracy : 0.8315
##
                    95% CI : (0.7373, 0.9025)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 1.018e-05
##
##
##
                     Kappa : 0.641
##
    Mcnemar's Test P-Value : 1
##
##
               Sensitivity: 0.7647
##
               Specificity: 0.8727
##
            Pos Pred Value: 0.7879
##
##
            Neg Pred Value: 0.8571
                Prevalence: 0.3820
##
            Detection Rate: 0.2921
##
      Detection Prevalence: 0.3708
##
##
         Balanced Accuracy: 0.8187
##
##
          'Positive' Class : Died
##
cat("\nF1: ", lda_f1, "\n")
##
## F1: 0.7761
lda probs <- predict(lda_model, newdata = test_data, type = "prob")</pre>
```

```
auc_value <- calculate_auc(test_data$DEATH_EVENT, lda_probs$Died) #Calculating AUC</pre>
cat("AUC:", auc_value, "\n\n\n")
```

```
# QDA
qda_pred <- predict(qda_model, newdata = test_data)</pre>
qda_cm <- confusionMatrix(qda_pred, test_data$DEATH_EVENT, positive = "Died")</pre>
qda_f1 <- calculate_f1(qda_cm)</pre>
print(qda_cm)
```

AUC: 0.9203

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    47
                         17
##
     Died
                     8
                         17
##
##
                  Accuracy : 0.7191
##
                    95% CI: (0.6138, 0.8093)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 0.02998
##
##
##
                     Kappa : 0.3734
##
    Mcnemar's Test P-Value: 0.10960
##
##
               Sensitivity: 0.5000
##
               Specificity: 0.8545
##
            Pos Pred Value: 0.6800
##
##
            Neg Pred Value : 0.7344
                Prevalence: 0.3820
##
            Detection Rate: 0.1910
##
      Detection Prevalence: 0.2809
##
##
         Balanced Accuracy: 0.6773
##
##
          'Positive' Class : Died
##
cat("\nF1: ", qda_f1, "\n")
##
## F1: 0.5763
qda_probs <- predict(qda_model, newdata = test_data, type = "prob")</pre>
```

```
auc_value <- calculate_auc(test_data$DEATH_EVENT, qda_probs$Died) #Calculating AUC</pre>
cat("AUC:", auc_value, "\n\n\n")
```

```
## AUC: 0.7289
```

```
# KNN
knn_pred <- predict(knn_model, newdata = test_data)</pre>
knn_cm <- confusionMatrix(knn_pred, test_data$DEATH_EVENT, positive = "Died")</pre>
knn_f1 <- calculate_f1(knn_cm)</pre>
print(knn_cm)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    48
##
     Died
                     7
                         15
##
##
                  Accuracy : 0.7079
##
                    95% CI: (0.6019, 0.7995)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 0.04905
##
##
##
                     Kappa: 0.3366
##
    Mcnemar's Test P-Value: 0.03098
##
##
               Sensitivity: 0.4412
##
               Specificity: 0.8727
##
            Pos Pred Value: 0.6818
##
##
            Neg Pred Value : 0.7164
                Prevalence: 0.3820
##
            Detection Rate: 0.1685
##
      Detection Prevalence : 0.2472
##
##
         Balanced Accuracy: 0.6570
##
##
          'Positive' Class : Died
##
cat("\nF1: ", knn_f1, "\n")
##
## F1: 0.5357
knn probs <- predict(knn_model, newdata = test_data, type = "prob")</pre>
```

```
knn_probs <- predict(knn_model, newdata = test_data, type = "prob")
auc_value <- calculate_auc(test_data$DEATH_EVENT, knn_probs$Died) #Calculating AUC
cat("AUC:", auc_value, "\n\n\n")</pre>
```

AUC: 0.8142

```
# SVM
svm_pred <- predict(svm_model, newdata = test_data)
svm_cm <- confusionMatrix(svm_pred, test_data$DEATH_EVENT, positive = "Died")
svm_f1 <- calculate_f1(svm_cm)
print(svm_cm)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    46
##
     Died
                     9
                         24
##
##
                  Accuracy : 0.7865
##
                    95% CI: (0.6869, 0.8663)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 0.000518
##
##
##
                     Kappa : 0.5453
##
    Mcnemar's Test P-Value : 1.000000
##
##
               Sensitivity: 0.7059
##
               Specificity: 0.8364
##
            Pos Pred Value: 0.7273
##
##
            Neg Pred Value : 0.8214
                Prevalence: 0.3820
##
            Detection Rate: 0.2697
##
      Detection Prevalence: 0.3708
##
##
         Balanced Accuracy: 0.7711
##
##
          'Positive' Class : Died
##
cat("\nF1: ", svm_f1, "\n")
##
## F1: 0.7164
svm probs <- predict(svm_model, newdata = test_data, type = "prob")</pre>
auc_value <- calculate_auc(test_data$DEATH_EVENT, svm_probs$Died) #Calculating AUC</pre>
```

```
cat("AUC:", auc_value, "\n\n\n")
## AUC: 0.8861
```

```
# Random Forest
rf_pred <- predict(rf_model, newdata = test_data)
rf_cm <- confusionMatrix(rf_pred, test_data$DEATH_EVENT, positive = "Died")
rf_f1 <- calculate_f1(rf_cm)
print(rf_cm)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    51
                         12
##
     Died
                     4
                         22
##
##
                  Accuracy : 0.8202
##
                    95% CI: (0.7245, 0.8936)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 3.019e-05
##
##
##
                     Kappa : 0.6013
##
    Mcnemar's Test P-Value: 0.08012
##
##
               Sensitivity: 0.6471
##
               Specificity: 0.9273
##
            Pos Pred Value: 0.8462
##
##
            Neg Pred Value: 0.8095
                Prevalence: 0.3820
##
            Detection Rate: 0.2472
##
      Detection Prevalence : 0.2921
##
##
         Balanced Accuracy: 0.7872
##
##
          'Positive' Class : Died
##
cat("\nF1: ", rf_f1, "\n")
##
## F1: 0.7333
rf_probs <- predict(rf_model, newdata = test_data, type = "prob")</pre>
```

```
auc_value <- calculate_auc(test_data$DEATH_EVENT, rf_probs$Died) #Calculating AUC</pre>
cat("AUC:", auc_value, "\n\n\n")
```

```
## AUC: 0.9516
```

```
# Decision Tree
dt_pred <- predict(dt_model, newdata = test_data)</pre>
dt_cm <- confusionMatrix(dt_pred, test_data$DEATH_EVENT, positive = "Died")</pre>
dt_f1 <- calculate_f1(dt_cm)</pre>
print(dt_cm)
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction Survived Died
##
     Survived
                    51
##
     Died
                     4
                         25
##
##
                  Accuracy : 0.8539
##
                    95% CI: (0.7632, 0.9199)
       No Information Rate: 0.618
##
       P-Value [Acc > NIR] : 9.176e-07
##
##
##
                     Kappa: 0.6817
##
    Mcnemar's Test P-Value: 0.2673
##
##
               Sensitivity: 0.7353
##
               Specificity: 0.9273
##
            Pos Pred Value: 0.8621
##
##
            Neg Pred Value: 0.8500
                Prevalence: 0.3820
##
            Detection Rate: 0.2809
##
      Detection Prevalence: 0.3258
##
##
         Balanced Accuracy: 0.8313
##
##
          'Positive' Class : Died
##
```

```
cat("\nF1: ", dt_f1, "\n")
```

```
##
## F1: 0.7937
```

```
dt_probs <- predict(dt_model, newdata = test_data, type = "prob")
auc_value <- calculate_auc(test_data$DEATH_EVENT, dt_probs$Died) #Calculating AUC
cat("AUC:", auc_value, "\n\n\n")</pre>
```

```
## AUC: 0.8639
```

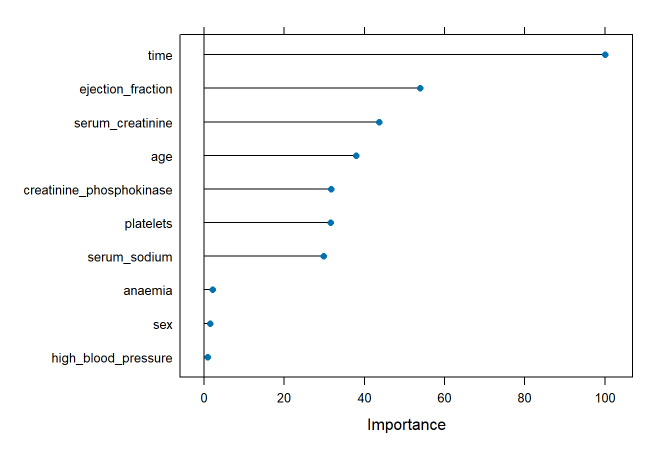
5.3 Variable Importance for Random Forest

```
# variable importance for rf
library(caret)
importance_rf <- varImp(rf_model)
print(importance_rf)</pre>
```

```
## rf variable importance
##
##
                             Overall
## time
                             100.0000
## ejection_fraction
                             53.9184
## serum_creatinine
                             43.7095
## age
                              37.9317
## creatinine_phosphokinase 31.6159
## platelets
                             31.5847
## serum_sodium
                             29.7679
## anaemia
                              2.1262
## sex
                               1.4597
## high_blood_pressure
                               0.9020
## smoking
                               0.6079
## diabetes
                               0.0000
```

```
# top 10 important variables
plot(importance_rf, top = 10, main = "Top 10 Important Variables - Random Forest")
```

Top 10 Important Variables - Random Forest



varImp() IDs which predictors had the biggest impact on model performance. In RF, var. importance is calculated based on how much each variable reduces impurity across all trees. The top variables are time, serum_creatine, ejection_fraction, and age, which aligns with domain knowledge-> indicators of heart failure progression.

6. Performance Comparision

6.1 Summary

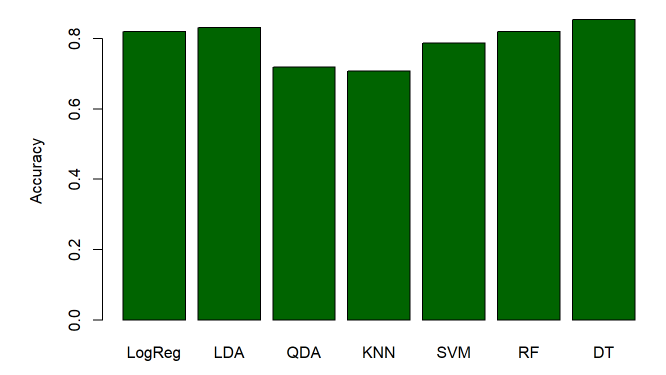
```
results <- resamples(list(SVM = svm_model, LogReg = logistic_regression_model, LDA = lda_model,
QDA = qda_model, RF = rf_model, KNN = knn_model))
summary(results)</pre>
```

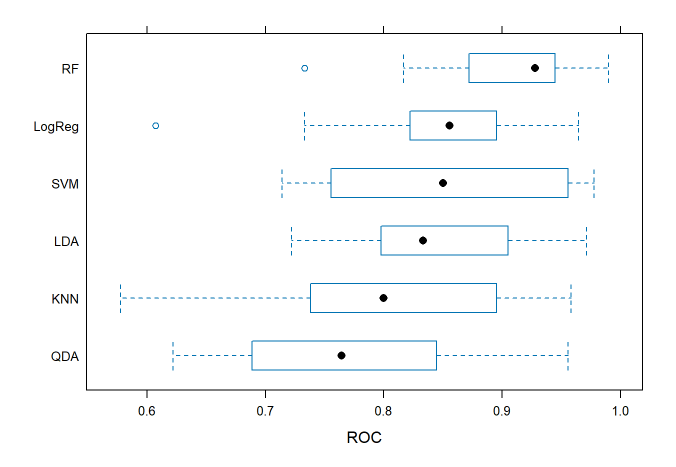
```
##
## Call:
## summary.resamples(object = results)
##
## Models: SVM, LogReg, LDA, QDA, RF, KNN
## Number of resamples: 10
##
## ROC
##
               Min.
                      1st Qu.
                                  Median
                                              Mean
                                                      3rd Qu.
                                                                   Max. NA's
## SVM
          0.7142857 0.7750000 0.8500000 0.8535714 0.9444444 0.9777778
## LogReg 0.6071429 0.8261905 0.8555556 0.8349206 0.8908730 0.9642857
                                                                            0
          0.7222222 0.8037698 0.8333333 0.8396032 0.8869048 0.9714286
## LDA
                                                                            0
## QDA
          0.6222222 0.7000000 0.7642857 0.7677778 0.8285714 0.9555556
                                                                            0
          0.7333333  0.8729167  0.9277778  0.8975907  0.9416667  0.9897959
## RF
                                                                            0
          0.5777778 0.7396825 0.8000000 0.8013889 0.8908730 0.9583333
                                                                            0
## KNN
##
## Sens
##
               Min.
                      1st Qu.
                                  Median
                                                      3rd Qu.
                                                                   Max. NA's
                                              Mean
          0.7142857 0.8821429 0.9333333 0.8842857 0.9333333 0.9333333
## SVM
## LogReg 0.7857143 0.8595238 0.9000000 0.8909524 0.9333333 1.0000000
                                                                            0
          0.8000000 0.8571429 0.8666667 0.8980952 0.9833333 1.00000000
## LDA
                                                                            0
## QDA
          0.7333333  0.8666667  0.8976190  0.8928571  0.9333333  1.0000000
                                                                            0
## RF
          0.8666667 0.9333333 0.9333333 0.9461905 0.9833333 1.0000000
                                                                            0
          0.8000000 0.9333333 0.9666667 0.9466667 1.0000000 1.0000000
## KNN
##
## Spec
##
               Min.
                      1st Qu.
                                  Median
                                              Mean
                                                      3rd Qu.
## SVM
          0.5000000 0.5000000 0.5833333 0.5928571 0.6666667 0.7142857
                                                                            0
## LogReg 0.3333333 0.5000000 0.6666667 0.6142857 0.6666667 1.0000000
                                                                            0
## LDA
          0.5000000 0.5416667 0.6666667 0.6404762 0.6666667 0.8571429
                                                                            0
## QDA
          0.1666667 0.3333333 0.3809524 0.4476190 0.6250000 0.8333333
                                                                            0
## RF
          0.3333333  0.4464286  0.5833333  0.5761905  0.6666667  1.0000000
                                                                            0
## KNN
          0.1428571 0.1666667 0.3333333 0.3380952 0.5000000 0.5714286
                                                                            0
```

6.2 Comparison Plots (Barplot and Boxplot of Model Performace)

```
# Boxplot for Accuracy
## bwplot(results, metric = "Accuracy")
get_accuracy <- function(cm) {</pre>
  accuracy <- as.numeric(cm$overall["Accuracy"])</pre>
  return(round(accuracy, 4))
}
logreg_acc <- get_accuracy(logreg_cm)</pre>
lda_acc <- get_accuracy(lda_cm)</pre>
qda_acc <- get_accuracy(qda_cm)</pre>
knn_acc <- get_accuracy(knn_cm)</pre>
svm_acc <- get_accuracy(svm_cm)</pre>
rf_acc <- get_accuracy(rf_cm)</pre>
dt_acc <- get_accuracy(dt_cm)</pre>
accuracy_df <- data.frame(Model = c("LogReg", "LDA", "QDA", "KNN", "SVM", "RF", "DT"),</pre>
                            Accuracy = c(logreg_acc, lda_acc, qda_acc, knn_acc, svm_acc, rf_acc, d
t_acc))
# Bar plot for accuracy - used this instead of boxplot because the accuracy were singe points so
boxplot was a line rather than box
barplot(accuracy_df$Accuracy, names.arg = accuracy_df$Model, col = "darkgreen",
        main = "Accuracy Comparison of Models", ylab = "Accuracy")
```

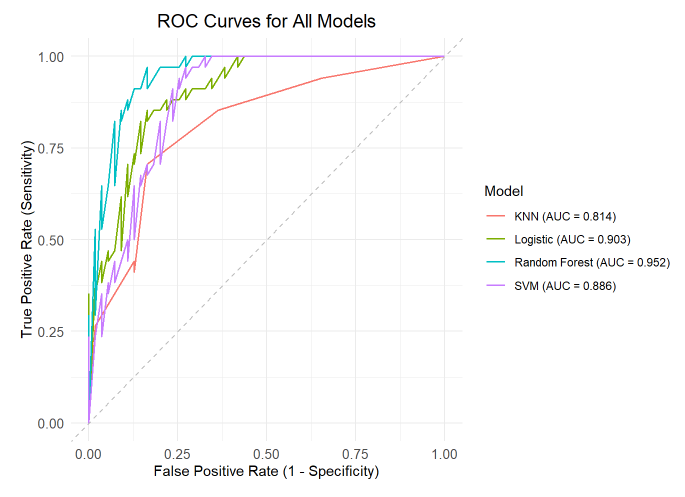
Accuracy Comparison of Models





6.3 ROC Curve

```
# function to extract ROC curve data
extract_roc_df <- function(roc_obj, model_name) {</pre>
 data.frame(
    FPR = 1 - roc_obj$specificities,
    TPR = roc_obj$sensitivities,
    Model = model_name
 )
}
true_labels <- test_data$DEATH_EVENT</pre>
# calculate ROC curves and AUCs
roc_logreg <- roc(true_labels, logreg_probs$Died)</pre>
         <- roc(true_labels, knn_probs$Died)</pre>
         <- roc(true_labels, svm_probs$Died)</pre>
roc svm
roc_rf
          <- roc(true_labels, rf_probs$Died)</pre>
# get AUC values
auc_logreg <- round(auc(roc_logreg), 3)</pre>
auc_knn
         <- round(auc(roc_knn), 3)</pre>
auc svm
         <- round(auc(roc svm), 3)</pre>
auc_rf
          <- round(auc(roc_rf), 3)</pre>
# combine data
roc_data <- rbind(</pre>
  extract_roc_df(roc_logreg, paste0("Logistic (AUC = ", auc_logreg, ")")),
 extract_roc_df(roc_knn, paste0("KNN (AUC = ", auc_knn, ")")),
 extract roc df(roc svm, paste0("SVM (AUC = ", auc svm, ")")),
  extract_roc_df(roc_rf, paste0("Random Forest (AUC = ", auc_rf, ")"))
)
# plot
ggplot(roc_data, aes(x = FPR, y = TPR, color = Model)) +
  geom\_line(size = 0.6) +
  geom_abline(slope = 1, intercept = 0, linetype = "dashed", color = "grey") +
  labs(title = "ROC Curves for All Models",
       x = "False Positive Rate (1 - Specificity)",
       y = "True Positive Rate (Sensitivity)") +
  theme_minimal() +
  theme(
    plot.title = element_text(hjust = 0.5, size = 14),
    axis.text = element_text(size = 10),
    axis.title = element_text(size = 11)
  )
```



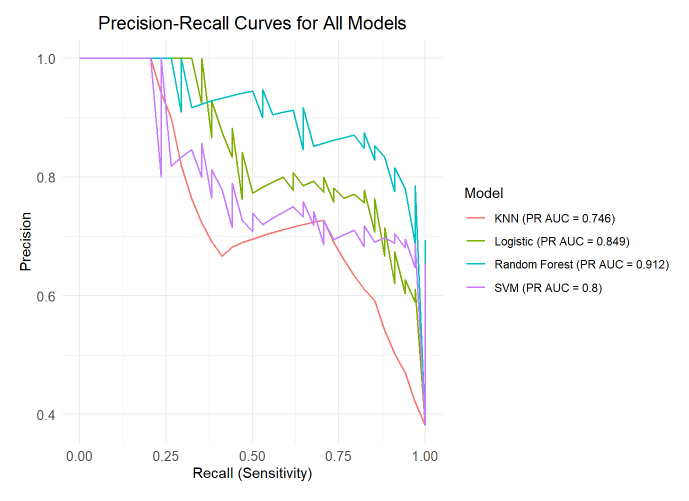
AUC shows probability that random positive class instance (Died) is ranked higher than random negative class (Survived) in predicted probability. It is more informative when class distribution is imbalanced (ex. hfp_data 68/32) in comparison to accuracy and F1-score since they are threshold-independent and not affected by skewed data.

- Random Forest performs the best (AUC=0.952), and the top-left curve indicates a high sensitive and low false positive rate
- Logistic Regression and Support Vector Machine perform strongly (AUC=0.903, 0.886 respectively)
- **K-Nearest Neighbour** is least effective out of the 4 (AUC=0.814). This might be because of noise, high variance, "curse of dimensionality".

6.4 PR Curves

```
# PR curves
pr_logreg <- pr.curve(scores.class0 = logreg_probs$Died[true_labels == "Died"],</pre>
                       scores.class1 = logreg_probs$Died[true_labels == "Survived"],
                       curve = TRUE)
pr_knn <- pr.curve(scores.class0 = knn_probs$Died[true_labels == "Died"],</pre>
                   scores.class1 = knn_probs$Died[true_labels == "Survived"],
                   curve = TRUE)
pr_svm <- pr.curve(scores.class0 = svm_probs$Died[true_labels == "Died"],</pre>
                   scores.class1 = svm_probs$Died[true_labels == "Survived"],
                   curve = TRUE)
pr_rf <- pr.curve(scores.class0 = rf_probs$Died[true_labels == "Died"],</pre>
                   scores.class1 = rf_probs$Died[true_labels == "Survived"],
                  curve = TRUE)
# extract PR curve as data frame
extract_pr_df <- function(pr_obj, model_name) {</pre>
 data.frame(
    Recall = pr obj$curve[, 1],
    Precision = pr_obj$curve[, 2],
    Model = model name
  )
}
# PR AUCs
auc pr logreg <- round(pr logreg$auc.integral, 3)</pre>
auc_pr_knn
            <- round(pr_knn$auc.integral, 3)</pre>
auc_pr_svm <- round(pr_svm$auc.integral, 3)</pre>
auc_pr_rf
             <- round(pr_rf$auc.integral, 3)</pre>
# combine to one df for ggplot
pr_data <- rbind(</pre>
  extract_pr_df(pr_logreg, paste0("Logistic (PR AUC = ", auc_pr_logreg, ")")),
 extract_pr_df(pr_knn, paste0("KNN (PR AUC = ", auc_pr_knn, ")")),
 extract_pr_df(pr_svm, paste0("SVM (PR AUC = ", auc_pr_svm, ")")),
  extract_pr_df(pr_rf, paste0("Random Forest (PR AUC = ", auc_pr_rf, ")"))
)
# plot
ggplot(pr_data, aes(x = Recall, y = Precision, color = Model)) +
  geom\_line(size = 0.6) +
  labs(title = "Precision-Recall Curves for All Models",
       x = "Recall (Sensitivity)",
       y = "Precision",
       color = "Model") +
 theme_minimal() +
  theme(
    plot.title = element_text(hjust = 0.5, size = 14),
    axis.text = element_text(size = 10),
```





PR curve focuses on positive class ("Died") by plotting precision vs. recall. More informative than ROC AUC when class distribution is imbalanced, since it directly evaluates model ability to identify true positives.

- Random Forest has highest AUC-PR (0.912) which shows its ability to identify actual deaths while also keeping high precision.
- Logistic Regression had strong performance at 0.849
- Support Vector Machine gives a moderate performance at 0.8
- **K-Nearest Neighbour** performed least well, at 0.746, showing it had more false positives and less effective in ID actual deaths.

7. Hyperparameter Tuning

7.1 K-Nearest Neighbours (KNN)

```
## k-Nearest Neighbors
##
## 210 samples
   12 predictor
##
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 189, 188, 189, 190, ...
## Resampling results across tuning parameters:
##
##
    k
        ROC
                   Sens
                              Spec
##
     3 0.7268821 0.9247619 0.3690476
##
     5 0.7342971 0.9390476 0.3190476
##
     7 0.7748129 0.9528571 0.2904762
##
     9 0.7773469 0.9661905 0.3761905
    11 0.7862812 0.9590476 0.3547619
##
##
    13 0.8033617 0.9657143 0.3380952
##
    15 0.8143537 0.9657143 0.3214286
##
    17 0.8136111 0.9657143 0.3047619
##
    19 0.8121769 0.9590476 0.3214286
##
    21 0.8132370 0.9590476 0.3071429
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was k = 15.
```

7.2 Support Vector Machine (SVM)

```
## Support Vector Machines with Radial Basis Function Kernel
##
## 210 samples
   12 predictor
##
##
    2 classes: 'Survived', 'Died'
##
## Pre-processing: centered (12), scaled (12)
## Resampling: Cross-Validated (10 fold)
  Summary of sample sizes: 189, 188, 189, 190, 189, 189, ...
  Resampling results across tuning parameters:
##
##
    C
          sigma ROC
                            Sens
                                       Spec
##
    0.25 0.25
                 0.7745238 0.8980952 0.31904762
##
    0.25 0.50
                 0.6031746 0.9523810
                                       0.03095238
##
    0.25 1.00
                 0.4580952 0.9800000 0.000000000
##
    0.25 2.00
                 0.4731746 0.9733333 0.00000000
    0.25 4.00
##
                 0.5439286 0.9933333 0.00000000
##
    0.50 0.25
                 0.7745238 0.9042857 0.35000000
##
    0.50 0.50
                 0.6876190 0.9457143 0.06190476
##
    0.50 1.00
                 0.3603175 0.9866667
                                       0.00000000
##
    0.50 2.00
                 0.3725397
                            0.9866667
                                       0.00000000
##
    0.50 4.00
                 0.4302381 0.9861905
                                      0.00000000
##
    1.00 0.25
                 0.7691270 0.9180952 0.31666667
##
    1.00 0.50
                 0.6876190 0.9447619 0.01428571
##
    1.00 1.00
                 0.4857143 0.9866667
                                       0.00000000
##
    1.00 2.00
                 0.4153968 0.9733333 0.01666667
    1.00 4.00
##
                 0.4203571 0.9861905 0.00000000
##
    2.00 0.25
                 0.7711111 0.9190476 0.28571429
##
    2.00 0.50
                 0.6808730 0.9519048 0.01666667
##
    2.00 1.00
                 0.4242857 0.9866667
                                       0.00000000
##
    2.00 2.00
                 0.4139683 0.9795238 0.00000000
    2.00 4.00
##
                 0.4557937 0.9928571 0.00000000
    4.00 0.25
##
                 0.7735714 0.8976190 0.23809524
##
    4.00 0.50
                 0.6282540 0.9590476 0.01428571
##
    4.00 1.00
                 0.4531746 0.9866667
                                       0.00000000
    4.00 2.00
##
                 0.3725397
                            0.9661905
                                       0.01666667
    4.00 4.00
##
                 0.4557937
                            0.9861905 0.00000000
##
## ROC was used to select the optimal model using the largest value.
## The final values used for the model were sigma = 0.25 and C = 0.25.
```

7.3 Random Forest (RF)

```
## Random Forest
##
## 210 samples
##
   12 predictor
    2 classes: 'Survived', 'Died'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 189, 189, 188, 189, 189, ...
## Resampling results across tuning parameters:
##
##
    mtry ROC
                     Sens
                                Spec
##
    2
           0.8855556 0.9528571 0.5809524
##
    3
           0.8860317 0.9395238 0.6261905
##
    4
          0.8883333 0.9052381 0.6261905
##
    5
          0.8854762 0.9057143 0.6261905
          0.8788889 0.8980952 0.5976190
##
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 4.
```

7.4 Decision Tree (DT)

```
## CART
##
## 210 samples
##
   12 predictor
##
    2 classes: 'Survived', 'Died'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
  Summary of sample sizes: 189, 189, 189, 189, 189, 188, ...
   Resampling results across tuning parameters:
##
##
           ROC
                      Sens
    ср
                                 Spec
##
    0.001 0.8065079
                      0.8790476
                                 0.6119048
##
    0.006
           0.8065079
                      0.8790476
                                 0.6119048
    0.011 0.8065079
##
                      0.8790476 0.6119048
##
    0.016 0.8065079 0.8790476 0.6119048
##
    0.021 0.8048413 0.8857143 0.6619048
##
    0.026 0.7987302 0.8857143 0.6452381
##
    0.031 0.7857143 0.8923810 0.6285714
##
    0.036 0.7593651 0.8990476 0.6285714
##
    0.041 0.7593651 0.8990476 0.6285714
##
    0.046 0.7593651 0.9057143 0.6285714
##
    0.051 0.7593651 0.9057143 0.6285714
##
    0.056 0.7638095 0.9323810 0.5952381
##
    0.061 0.7638095 0.9323810 0.5952381
##
    0.066 0.7638095 0.9323810 0.5952381
##
    0.071 0.7638095 0.9323810 0.5952381
##
    0.076 0.7638095 0.9323810 0.5952381
    0.081 0.7638095 0.9323810 0.5952381
##
##
    0.086 0.7638095 0.9323810 0.5952381
##
    0.091 0.7638095 0.9323810 0.5952381
##
    0.096 0.7638095
                      0.9323810 0.5952381
##
## ROC was used to select the optimal model using the largest value.
## The final value used for the model was cp = 0.016.
```

For each model trained, we performed grid search and 10-fold cross validation to optimize k, c/sigma, mtry, and cp for their respective models. ROC was used to select optimal model using largest value.

8. Principal Component Analysis (PCA)

We applied PCA to reduce dimensionality so we can plot and interpret better, as PCA will give us a low-dimensional representation of the data that captures as much variance as possible.

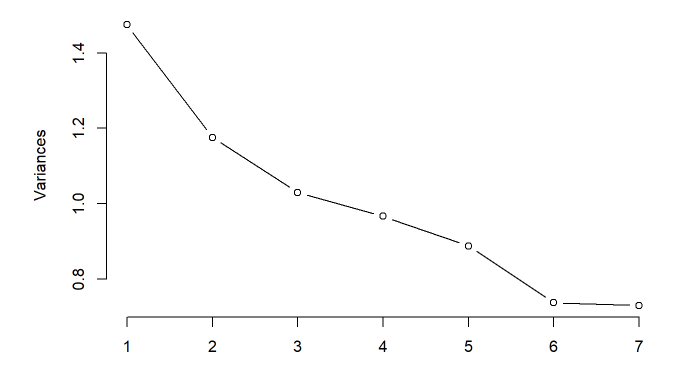
```
# extract only scaled continuous predictors
pca_data <- hfp_scaled[, continuous_vars]
pca_result <- prcomp(pca_data, center = FALSE, scale. = FALSE)</pre>
```

8.1 Scree Plot

This plot shows how much variance is explained by each principal component.

```
# scree plot -> proportion of variance explained
screeplot(pca_result, type = "lines", main = "Scree Plot of Principal Components")
```

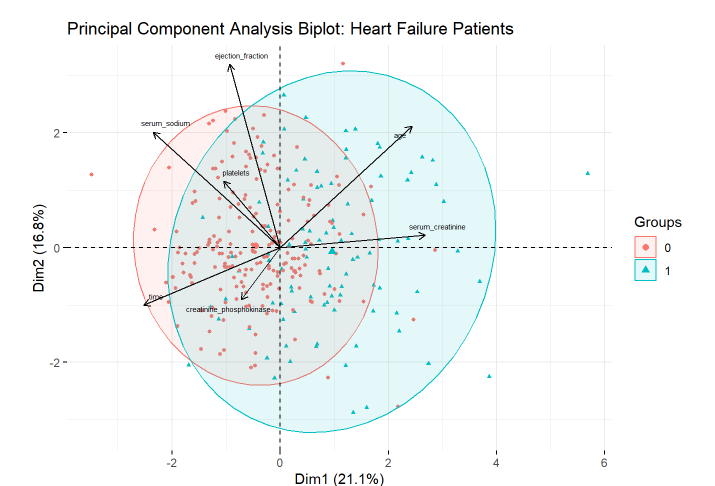
Scree Plot of Principal Components



scree plot: shows amt of variance explained by each PC, each point represents 1 PC and y-axis shows how much variance PC explains. We used it to determine how many PCs show meaningful dimensionality reduction - PC1 and PC2 explain most variance.

8.2 PCA Biplot

This plot shows patient scores on the first two PCs and the loadings of each variable.



This biplot visualizes both PCs (Dim1, Dim2) for each patient, and variable loadings. A positive loading shows that the variable increases the PC score, and a negative loading shows that the variable decreases the PC score. Each arrow represents a variable and coneys:

- · direction: how variable contributes to PCs
- length: how strongly a variable influences PCA (longer=stronger)
- · angle between arrows:
 - same direction = **positive correlation** between variables
 - opposite direction = **negative correlation** between variables
 - perpendicular = uncorrelated

This helps us visually see the clutering between survived (0) and died (1).

We can see that variables like serum_creatine and age arrows are **long and point right**, therfore they strongly contribute to PC1.

ejection_fraction and serum_sodium point towards the **upper-left** quadrant of the plot, and it has moderate positive loading on PC2, but negative on PC1.

time points bottom-left, meaning it loads negatively on PC1 and PC2.

creatine_phosphokinase points downward, and it contributes mostly negatively to PC2

platlets points left, indicating a negative PC1 loading

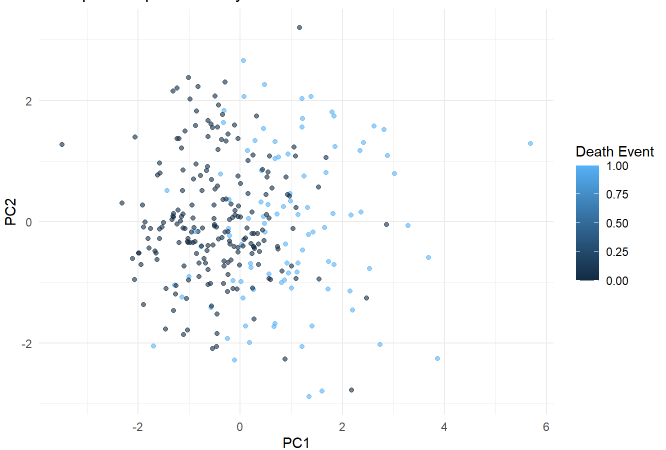
8.3 PC1 vs. PC2 Scatterplot

This plot visualizes patients in the PCA space, colored by survival outcome.

```
# PCA dataframe
pca_df <- as.data.frame(pca_result$x)
pca_df$DEATH_EVENT <- hfp_scaled$DEATH_EVENT

# PC1 vs. PC2
ggplot(pca_df, aes(x = PC1, y = PC2, color = DEATH_EVENT)) +
    geom_point(alpha = 0.6) +
    labs(title = "Principal Component Analysis: PC1 vs. PC2", x = "PC1", y = "PC2", color = "Death Event") +
    theme_minimal()</pre>
```

Principal Component Analysis: PC1 vs. PC2



The scatterplot explores how individual points lie in PCA space. It shows the spatial grouping of DEATH_EVENT classes, but it is not perfectly seperable. It also shows that PCA doesn't fully separate classes but shows patterns instead.

9. K-Means Clustering

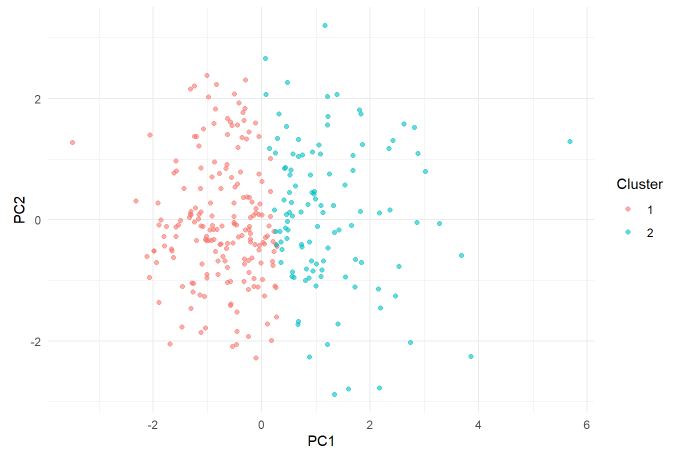
We applied K-Means (k=2) to identify groups in PCA reduced space, this will explore **unsupervised** grouping.

```
set.seed(2025)
# top 2 PCs for visualization, full data for clustering
kmeans_result <- kmeans(pca_data, centers = 2, nstart = 25)
# cluster labels to PCA df
pca_df$Cluster <- as.factor(kmeans_result$cluster)
# confusion matrix to check alignment with DEATH_EVENT
table(Cluster = pca_df$Cluster, DEATH_EVENT = pca_df$DEATH_EVENT)</pre>
```

```
## DEATH_EVENT
## Cluster 0 1
## 1 162 26
## 2 41 70
```

```
# plot clusters
ggplot(pca_df, aes(x = PC1, y = PC2, color = Cluster)) +
  geom_point(alpha = 0.6) +
  labs(title = "K-Means Clustering (k = 2) on PCA Components") +
  theme_minimal()
```

K-Means Clustering (k = 2) on PCA Components



We used k-means (k=2) on first 2 PC to see the unsupervised grouping of patients. Compared to the other scatter plot above which used <code>DEATH_EVENT</code>, K-Means formed the groups based only on patterns in the data and no knowledge of survival outcomes. The plot shows two distinct clusters, which indicates the dataset has a natural structure, and the patients are grouped based on similar features (ex. <code>age</code>, <code>serum_creatine</code>, etc) which had a strong impact on patterns found by PCA (plots above).