Conditional Inference Trees & Cox Regression to Predict Heart Failure Survival Time

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https://archive.ics.uci.edu/ml/datasets/Heart+failure+clinical+record/ (https://archive.ics.uci.edu/ml/datasets/Heart+failure+clinical+record/

• All 299 patients had left ventricular systolic dysfunction

Initial Variables:

- age: age of the patient (years)
- anaemia: presence of critically low haematocrit levels (boolean)
- high blood pressure: if the patient has hypertension (boolean)
- creatinine phosphokinase (CPK): level of the CPK enzyme in the blood (mcg/L)
- diabetes: if the patient has diabetes (boolean)
- ejection fraction: percentage of blood leaving the heart at each contraction (percentage)
- · platelets: platelets in the blood (kiloplatelets/mL)
- sex: woman or man (binary)
- . serum creatinine: level of serum creatinine in the blood (mg/dL)
- serum sodium: level of serum sodium in the blood (mEq/L)
- smoking: if the patient smokes or not (boolean)
- · time: follow-up period (days)
- [target] death event: if the patient deceased during the follow-up period (boolean)

```
library(ggplot2)
library(dplyr)
library(tidyr)
library(survival)
library(survminer)
library(partykit)
library(coin)
library(survminer)
library(flexsurv)
library(flexsurv)
library(randomForestSRC)
library(broom)
library(gstsummary)
```

Loading in the data

Creating Left Ventricular Ejection Fraction Groups set by Cardiology Experts (https://www.ncbi.nlm.nih.gov/books/NBK459131/). Rounding for averages instead of only using data for men and women.

```
HF <- read.csv("heart_failure_clinical_records_dataset.csv")

HF$anaemia = as.factor(HF$anaemia)
HF$diabetes = factor(HF$diabetes,levels=c(0,1),labels=c("Absent","Present"))
HF$hypertension = factor(HF$high_blood_pressure,levels=c(0,1),labels=c("Absent","Present"))

HF$sex = factor(HF$sex,levels=c(0,1),labels=c("Female","Male"))
HF$smoking = as.factor(HF$smoking)
HF$DEATH_EVENT = as.factor(HF$DEATH_EVENT)</pre>

HF <- select(HF, -high_blood_pressure)

skim(HF)
```

Data summary

Name	HF
Number of rows	299
Number of columns	13

Column type frequency:	
factor	6
numeric	7

Variable type: factor

skim_variable	n_missing	complete_rate ordered	n_unique top_counts
anaemia	0	1 FALSE	2 0: 170, 1: 129
diabetes	0	1 FALSE	2 Abs: 174, Pre: 125
sex	0	1 FALSE	2 Mal: 194, Fem: 105
smoking	0	1 FALSE	2 0: 203, 1: 96
DEATH_EVENT	0	1 FALSE	2 0: 203, 1: 96
hypertension	0	1 FALSE	2 Abs: 194, Pre: 105

Variable type: numeric

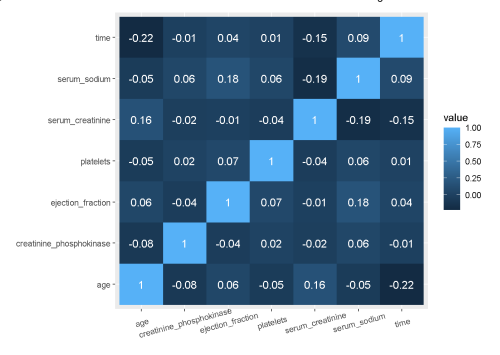
skim_variable	n_missing	complete_rate	mean	sd	p0	p25	p50	p75	p100	hist
age	0	1	60.83	11.89	40.0	51.0	60.0	70.0	95.0	
creatinine_phosphokinase	0	1	581.84	970.29	23.0	116.5	250.0	582.0	7861.0	-
ejection_fraction	0	1	38.08	11.83	14.0	30.0	38.0	45.0	80.0	_=
platelets	0	1	263358.03	97804.24	25100.0	212500.0	262000.0	303500.0	850000.0	_=_
serum_creatinine	0	1	1.39	1.03	0.5	0.9	1.1	1.4	9.4	■
serum_sodium	0	1	136.63	4.41	113.0	134.0	137.0	140.0	148.0	
time	0	1	130.26	77.61	4.0	73.0	115.0	203.0	285.0	

Correlation

Time and Serum_Creatinine have a correlation to Serum_Sodium of 0.15 & 0.19, respectively.

```
cormat <- HF %>% select(where(is.numeric)) %>% cor() %>% round(2)
melted_cormat <- reshape2::melt(cormat)

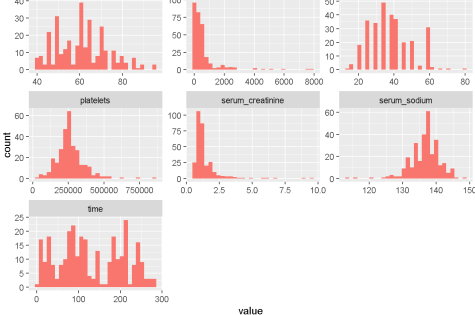
ggplot(data = melted_cormat, aes(x=Var1, y=Var2, fill=value)) +
    geom_tile() +
    geom_text(aes(Var2, Var1, label = value), color = "white", size = 4) +
    theme(axis.title.x=element_blank(),
        axis.title.y=element_blank(),
        axis.title.y=element_text(angle = 15, vjust = 0.8)
    )
</pre>
```



Choosing to grab distributions based on having hypertension- what's traditionally seen as a good indicator of heart failure.

Doing so to look at, specifically, Ejection Fraction right after to see if there is correlation.

```
HF %>%
 purrr::keep(is.numeric) %>%
  gather() %>%
 ggplot(aes(value)) +
    facet_wrap(~ key, scales = "free") +
    geom_histogram(aes(fill="orange"), show.legend = FALSE)
                                                                                 ejection_fraction
                                           creatinine_phosphokinase
                                    100
  40
                                                                     50
                                                                     40 -
  30
                                    75
                                                                     30
  20
                                    50
```

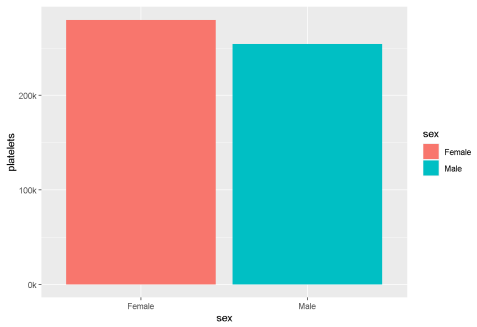


Comparing creatinine_phosphokinase to Men & Women— those who smoke and those who do not.

• Noticing that the average creatinine_phosphokinase is higher for non-smokers.

```
ggplot(HF, aes(x=sex, y=platelets, fill=sex)) +
  geom_bar(position = "dodge", stat="summary", fun="mean") +
  scale_y_continuous(labels = scales::label_number(suffix = "k", scale = 1e-3)) +
  ggtitle("Mean Platelets for Sexes")
```





```
HF %>% group_by(sex, DEATH_EVENT) %>%
summarize(count = n(), .groups="drop")
```

```
## # A tibble: 4 × 3

## sex DEATH_EVENT count

## <fct> <fct> <int>
## 1 Female 0 71

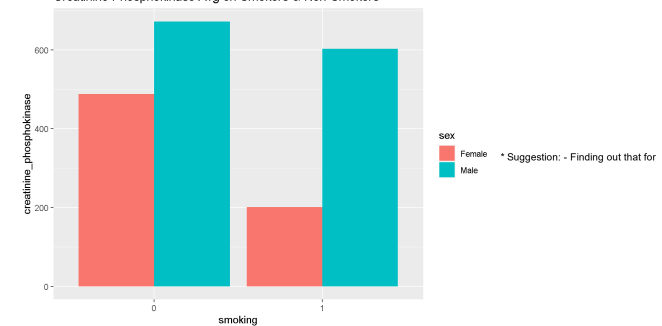
## 2 Female 1 34

## 3 Male 0 132

## 4 Male 1 62
```

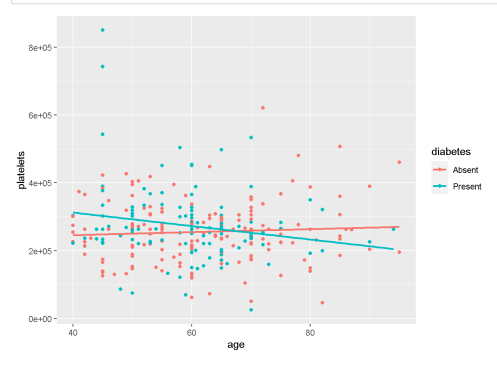
```
ggplot(HF, aes(x=smoking, y=creatinine_phosphokinase, fill=sex)) +
geom_bar(position = "dodge", stat="summary", fun="mean") +
ggtitle("Creatinine Phosphokinase Avg on Smokers & Non-Smokers")
```

Creatinine Phosphokinase Avg on Smokers & Non-Smokers



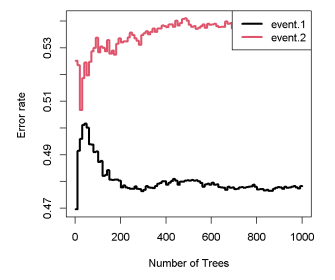
those diabetic, plateletes reduce as age increases. - For those who aren't diabetic, plateletes generally stay the same and potentially, increase by a marginal amount.

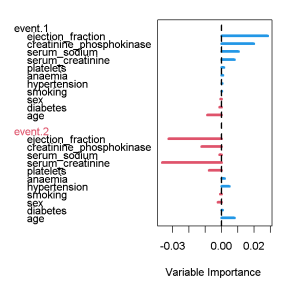
```
ggplot(HF, aes(x=age, y=platelets,color=diabetes)) + geom_point() +
  geom_smooth(method='lm', se = FALSE)
```



Random Forest Survival

Used to get variable importance chart.



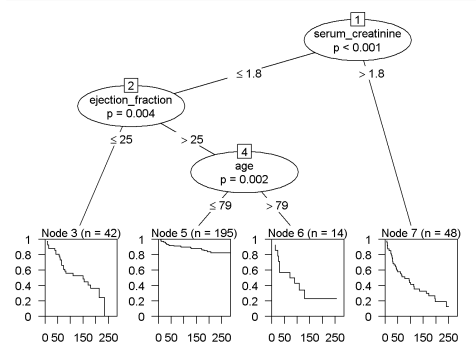


```
##
                               event.1
                                         event.2
## ejection_fraction
                               0.0283
                                         -0.0326
## creatinine phosphokinase
                               0.0197
                                         -0.0124
                               0.0103
                                         -0.0013
## serum sodium
## serum_creatinine
                               0.0078
                                         -0.0366
## platelets
                                         -0.0077
                               0.0014
## anaemia
                               0.0009
                                         0.0019
## hypertension
                               0.0003
                                          0.0047
## smoking
                                0.0002
                                         -0.0008
## sex
                               -0.0007
                                         -0.0023
## diabetes
                               -0.0013
                                          0.0005
```

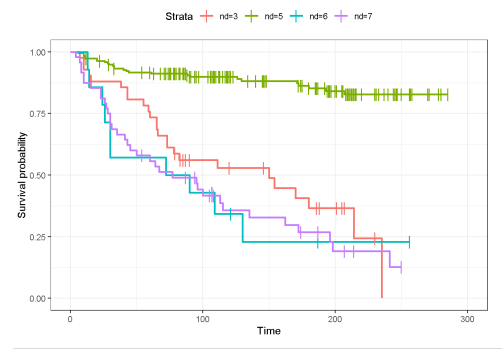
Conditional Inference Trees - Kaplan Meier Curves

We can see we have remaining cases in which the person did was not declared deceased due to the ending of the curve not dropping down to n%

Insights from this graph include: * Serum Creatinine is highly significant with the showcased split at 1.8 for survival prediction.



Plotting all node distributions/curves in one plot.



```
# Extracting survival curve for only one observation from the ctree. Perhaps an outlier.
#nd1 <- predict(CondInfTree, type = "prob")[[10]]
#summary(nd1, times=c(20, 45, 60, 80, 100, 10*(11:15)))
```

Constructing an exponential curve for previous graph's second node. * 48% probability of survival after t=150 days for patients older that have less than or equal to 1.8 in serum creatine, and an ejection fraction under 25.

```
K <- HF %>%
  filter(serum_creatinine <= 1.8, ejection_fraction <= 25)

# This one is best.
# The ~ 1 is our way ofletting R know that we aren't using any x variables. Just time and whether event occured which are both y variabes.
pred_k_surv <- survfit(Surv(time, DEATH_EVENT) ~ 1, data = K)
summary(pred_k_surv, times=c(20, 45, 60, 80, 100, 10*(11:15)))</pre>
```

```
## Call: survfit(formula = Surv(time, DEATH_EVENT) ~ 1, data = K)
##
##
   time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
     20
            36
                     5
                          0.881 0.0500
                                               0.788
                                                            0.985
##
                                                            0.937
      45
            33
                     3
                          0.808 0.0612
                                               0.696
##
     60
            31
                     3
                          0.734 0.0688
                                               0.611
                                                            0.882
                                                            0.759
##
     80
            23
                     6
                          0.587 0.0768
                                               0.454
##
    100
            17
                          0.562 0.0776
                                               0.429
                                                            0.736
                     1
##
    110
            17
                     0
                          0.562 0.0776
                                               0.429
                                                            0.736
##
    120
                          0.529 0.0798
                                               0.393
                                                            0.711
##
    130
            14
                     0
                          0.529 0.0798
                                               0.393
                                                            0.711
##
                     a
    140
            14
                          0.529 0.0798
                                               0.393
                                                            0.711
##
    150
            13
                     1
                          0.488
                                 0.0834
                                               0.349
                                                            0.682
```

- No pruning was done since most trees found revolve around the same 3 variables.
- Probability of survival after 150 days for those younger than 70 is 77%.
- Probability of survival after 200 days for those younger than 70 is 70%.

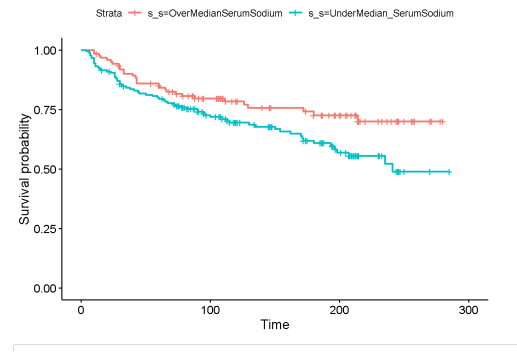
```
survfit(Surv(time, DEATH_EVENT) ~ 1, data = HF %>% filter(age <= 70)) %>%
tbl_survfit(
  times = c(150,200),
  label_header = "**{time} Day Survival (95% CI) For Those Younger Than 70**"
)
```

Characteristic	150 Day Survival (95% CI) For Those Younger Than 70	200 Day Survival (95% CI) For Those Younger Than 70				
Overall	77% (71%, 82%)	70% (64%, 77%)				

Looking at Serum Sodium Splitting at the median in case this dataset has any bias bc of outliers. * Finding that those with higher serum sodium have better survival rates, on average.

```
survfit(Surv(time, DEATH_EVENT) ~ 1, data = HF %>% filter(creatinine_phosphokinase <= 70)) %>%
tbl_survfit(
  times = c(150,200),
  label_header = "**{time} Day Survival (95% CI) For Those Younger Than 70**"
)
```

Characteristic	150 Day Survival (95% CI) For Those Younger Than 70	200 Day Survival (95% CI) For Those Younger Than 70		
Overall 72% (57%, 90%)		72% (57%, 90%)		
ss <- HF %>% mutate(s_s = i	felse((serum_sodium <= median(serum_sodium)), "UnderMed	ian_SerumSodium", "OverMedianSerumSodium"))		
ss_fit <- survfi	t(Surv(time, DEATH_EVENT) ~ s_s, data=ss)			
ggsurvnlot(ss fi	t data = ss)			



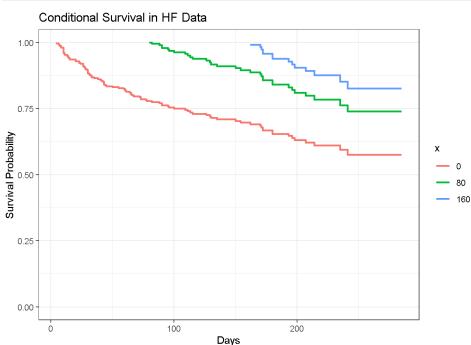
#CLEARLY, HAVING HIGHER SERUM_SODIUM MEANS HIGHER RATE OF SURVIVAL.

Conditional Survival: The probability of surviving further 't' years, given that a patient has already survived 's' years

```
remotes::install_github("zabore/condsurv")
library(condsurv)

fit1 <- survfit(Surv(time, DEATH_EVENT) ~ 1, data = HF)

gg_conditional_surv(
   basekm = fit1,
   at = seq(0, 160, 80),
   main = "Conditional Survival in HF Data",
   xlab = "Days",
   ylab = "Survival Probability"
   )</pre>
```



Cox Proportional Hazards Model (Cox Regression)

KM will make the curve based on event & time but that's all. We need to include the rest of the variables.

- At a given instance in time, someone who has hypertension is 0.42 times as likely to die as someone without hypertension adjusting for age.
- At any given instance in time, someone who does not have hypertension is 0.65 times as likely to die as someone who does, adjusting for age.
- · Concordance: Goodness of fit for survival analysis.

```
# hypertension useful bc tree didn't output it. i paired it w/ age bc why not?
coxMod2 <- coxph(Surv(time, DEATH_EVENT) ~ hypertension + age, data=HF)
summary(coxMod2)</pre>
```

```
## coxph(formula = Surv(time, DEATH_EVENT) ~ hypertension + age,
##
      data = HF)
##
##
    n= 299, number of events= 96
##
##
                         coef exp(coef) se(coef)
                                                   z Pr(>|z|)
## hypertensionPresent 0.417717 1.518491 0.209708 1.992 0.0464 *
## age
                     0.042424 1.043336 0.008693 4.880 1.06e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
                     exp(coef) exp(-coef) lower .95 upper .95
## hypertensionPresent
                         1.518
                                  0.6585
                                            1.007
## age
                         1.043
                                  0.9585
                                             1.026
                                                      1.061
##
## Concordance= 0.638 (se = 0.031 )
## Likelihood ratio test= 27.36 on 2 df, p=1e-06
## Wald test = 27.52 on 2 df, p=1e-06
## Score (logrank) test = 28.25 on 2 df, p=7e-07
```

Checking Linearity of Model * Linearity of the final cox regression is sufficient. * Anaemia is not statistically significant.

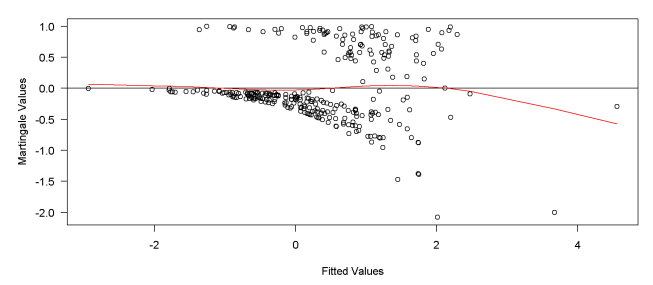
```
## Call:
## coxph(formula = Surv(time, DEATH EVENT) ~ age + anaemia + creatinine phosphokinase +
      ejection fraction + serum creatinine + hypertension, data = HF)
##
##
   n= 299, number of events= 96
##
##
                              coef exp(coef) se(coef)
                                                           z Pr(>|z|)
                          4.361e-02 1.045e+00 8.853e-03 4.926 8.41e-07 ***
## age
## anaemia1
                          3.933e-01 1.482e+00 2.129e-01 1.847 0.0648 .
## creatinine_phosphokinase 1.965e-04 1.000e+00 9.856e-05 1.993 0.0462 *
## ejection_fraction -5.179e-02 9.495e-01 1.005e-02 -5.152 2.57e-07 ***
## serum_creatinine
                         3.483e-01 1.417e+00 6.550e-02 5.318 1.05e-07 ***
## hypertensionPresent 4.668e-01 1.595e+00 2.129e-01 2.192 0.0284 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
                         exp(coef) exp(-coef) lower .95 upper .95
## age
                           1.0446 0.9573 1.0266 1.0629
                                      0.6749
                                                       2.2493
## anaemia1
                           1.4818
                                              0.9762
## creatinine_phosphokinase 1.0002
                                     0.9998
                                              1.0000
                                                        1.0004
## ejection_fraction
                           0.9495
                                     1.0531
                                               0.9310
                                                        0.9684
## serum_creatinine
                           1.4167
                                      0.7059
                                              1.2460
                                                        1.6108
## hypertensionPresent
                           1.5948
                                      0.6270 1.0506
                                                       2,4209
## Concordance= 0.738 (se = 0.028)
## Likelihood ratio test= 77.02 on 6 df, p=1e-14
## Wald test
                    = 85.82 on 6 df, p=2e-16
## Score (logrank) test = 83.51 on 6 df, p=7e-16
```

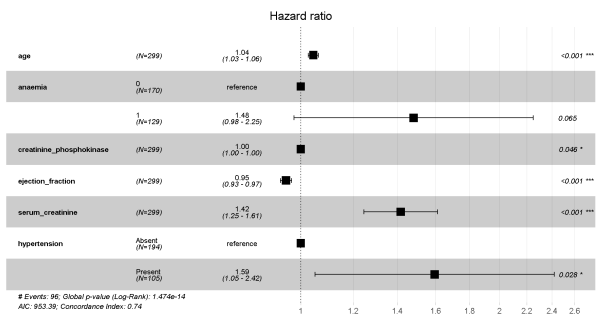
```
plot(predict(sigMod), residuals(sigMod, type = "martingale"), xlab = "Fitted Values",
   ylab = "Martingale Values", main = "Residual Plot", las = 1) +
   abline(h=0) +
   lines(smooth.spline(predict(sigMod), residuals(sigMod, type="martingale")), col="red")
```

```
## integer(0)
```

```
ggforest(sigMod, data = HF)
```

Residual Plot

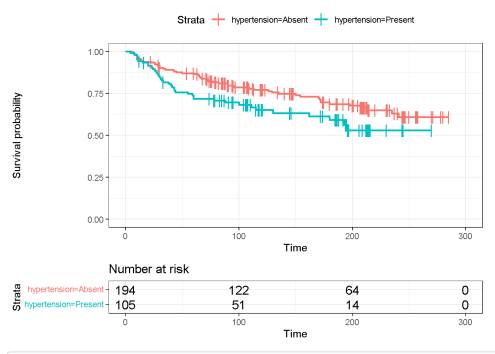




```
library(ggthemes)
finMod <- sigMod %>% tidy()
# finMod %>% mutate(upper = estimate + 1.96 * std.error,
           lower = estimate - 1.96 * std.error) %>%
   mutate(across(all_of(c("estimate", "lower", "upper")), exp)) %>%
    ggplot(aes(estimate, term, color = estimate > 1)) +
    geom_vline(xintercept = 1, color = "gray75") +
    geom linerange(aes(xmin = lower, xmax = upper), size = 2.25, alpha = 0.28) +
    geom_point(size = 4) +
    theme\_gdocs(base\_size = 16) +
    scale_color_manual(values = c("green4", "red3"), guide = "none") +
    x \lim(c(0, 3)) +
    labs(title = "Hazard Ratios for Significant Variables", y = NULL,
#
        x = "Hazard Ratio Estimates (95% C.I.)") +
#
    theme(axis.text.y = element_text(hjust = 0, size = 18)) +
#
      geom_text(label = exp(finMod$estimate) %>% round(2),
                nudge_y = .2, nudge_x = .15)
```

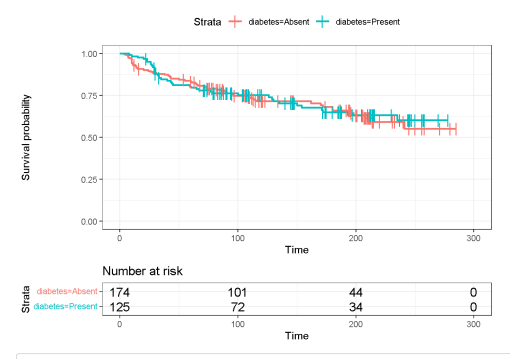
Performing the Log-Rank Test on select variables to extract significance.

.



```
survdiff(Surv(time,DEATH_EVENT) ~ hypertension, data=HF)
```

```
## Call:
## survdiff(formula = Surv(time, DEATH_EVENT) ~ hypertension, data = HF)
##
##
                         N Observed Expected (0-E)^2/E (0-E)^2/V
## hypertension=Absent 194
                                 57
                                        66.4
                                                  1.34
                                                            4.41
## hypertension=Present 105
                                 39
                                        29.6
                                                   3.00
                                                            4.41
##
   Chisq= 4.4 on 1 degrees of freedom, p= 0.04
```



```
## Call:
## call:
## survdiff(formula = Surv(time, DEATH_EVENT) ~ diabetes, data = HF)
##
##
##

N Observed Expected (0-E)^2/E (0-E)^2/V
## diabetes=Absent 174 56 55 0.0172 0.0405
## diabetes=Present 125 40 41 0.0231 0.0405
##
## Chisq= 0 on 1 degrees of freedom, p= 0.8
```

Binary Logistic Regression

Creating category variables for Serum Creatinine & Creatinine Phosphokinase due to their heavy right skewness.

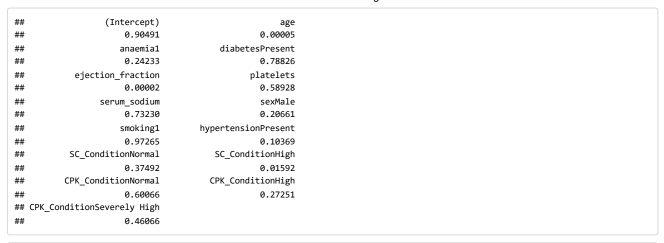
 $Using One (https://labs.selfdecode.com/blog/creatine-kinase/\#:\sim:text=The\%20low\%20normal\%20limit\%20for, 3\%2C\%204\%2C\%205\%5D./) \ and Two (https://www.mayoclinic.org/tests-procedures/creatinine-test/about/pac-$

20384646#:~:text=The%20typical%20range%20for%20serum,52.2%20to%2091.9%20micromoles%2FL).

```
set.seed(0)
library(caTools)

split_log <- sample.split(HF, SplitRatio = 0.7)
train_log <- subset(HF, split_log == TRUE) %>% select(-time)
test_log <- subset(HF, split_log == FALSE)

logit1 <- glm(DEATH_EVENT~., family = binomial,data = train_log)
summary(logit1)$coefficients[,4] %>% round(digits = 5)
```



summary(logit1)\$aic

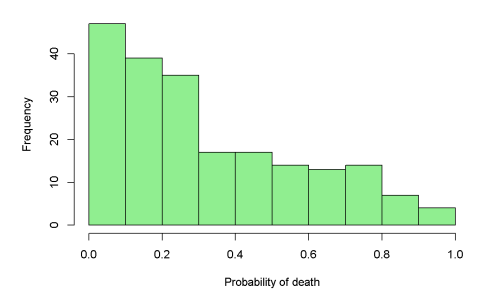
[1] 217.2686

```
logit2 <- step(logit1, direction = "backward", trace = FALSE)
summary(logit2)$coefficients[,4] %>% round(digits = 5)
```

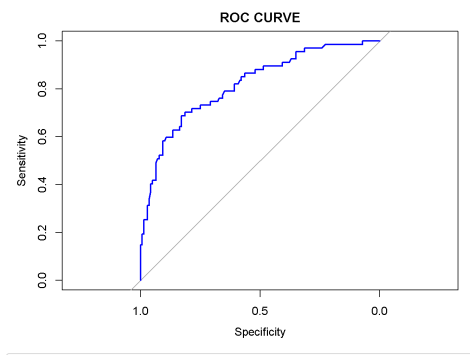
```
## (Intercept) age ejection_fraction hypertensionPresent
## 0.00349 0.00003 0.00010 0.04829
## SC_ConditionNormal SC_ConditionHigh
## 0.40556 0.01190
```

hist(logit2\$fitted.values, main=" Histogram ",xlab="Probability of death", col='light green')

Histogram



r <- pROC::roc(DEATH_EVENT~logit2\$fitted.values, data = train_log, plot = TRUE, main = "ROC CURVE", col= "blue")



```
optimal_roc <- r$thresholds[which.max(r$sensitivities + r$specificities)]

test_predicted_data <- test_log %>%
    mutate(p1=predict(logit2, newdata=test_log, type="response")) %>%
        mutate(Predict=ifelse(p1 < optimal_roc,0,1))

cm <- table(test_predicted_data$DEATH_EVENT,test_predicted_data$Predict) %>% prop.table()
    rownames(cm) <- c("Obs. neg","Obs. pos")
    colnames(cm) <- c("Pred. neg","Pred. pos")

ERROR.RESULTS <- tibble(
    Sensitivity=c(cm[1,1]/sum(cm[1,])),
    Specificity=c(cm[2,2]/sum(cm[2,])),
    FalsePositives=c(cm[2,2]/sum(cm[2,])),
    FalseNegatives=c(cm[1,2]/sum(cm[1,]))
)

efficiency <- sum(diag(cm))/sum(cm)

ERROR.RESULTS</pre>
```

```
## # A tibble: 1 × 4
## Sensitivity Specificity FalsePositives FalseNegatives
## <dbl> <dbl> <dbl> <dbl> <dbl> 
## 1 0.698 0.586 0.414 0.302
```

```
efficiency
```

```
## [1] 0.6630435
```

Findings:

- At a given instance in time, someone who has hypertension is 0.42 times as likely to die as someone without hypertension adjusting for age.
- Probability of survival after 150 days for those younger than 70 is 77%.
- Probability of survival after 200 days for those younger than 70 is 70%.
- 24% probability of survival after t=130 days for patients older than 79, that have less than or equal to 1.8 in serum creatine, and an ejection fraction over 25.

- For those diabetic, plateletes seem to reduce as age increases.
- On average, creatinine_phosphokinase is higher for non-smokers.
- Men, on average, have higher creatinine_phosphokinase.
- Women, on average, have a higher platelets count.
- age, ejection fraction, the presence of hypertension, and a value of serum creatinine greater than 1.25 are the variables that contribute most to an accurate prediction of mortality.
- age, creatinine_phosphokinase, ejection_fraction, serum_creatinine, and the presence of hypertension are what most impact the survival rate probability.
- anemia, smoking, sex status, and diabetes, and are the fields that contribute the least to survival rates, in that order.