## MVA Assignment 3

Tanvi

10/01/2020

### Assignment 3 - Data Cleaning, EDA, Tests

This document does a preliminary analysis on the Heart Failure Prediction dataset

### We will start by loading libraries and data

# clear environment rm(list = ls()) # defining libraries library(ggplot2) library(dplyr) library(PerformanceAnalytics) library(data.table) library(sqldf) library(nortest) library(tidyverse) 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)

# reading data

data <- read.csv('/Users/mac/Downloads/heart\_failure\_clinical\_records\_dataset.csv') # structure of data
str(data)</pre>

## '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...
                                        : num 265000 263358 162000 210000 327000 ...
## $ platelets
                                        : num 1.9 1.1 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.4 ...
## $ serum_creatinine
                                        : int 130 136 129 137 116 132 137 131 138 133 ...
##$ serum sodium
## $ sex
                                        : int 1 1 1 1 0 1 1 1 0 1 ...
                                        : int 0 0 1 0 0 1 0 1 0 1 ...
## $ smoking
## $ time
                                        : int 4 6 7 7 8 8 10 10 10 10 ...
## $ DEATH EVENT
                                        : int 1 1 1 1 1 1 1 1 1 1 ...
glimpse(data)
## Observations: 299 ##
Variables: 13
## $ age <dbl> 75, 55, 65, 50, 65, 90, 75, 60, 65, 8... ## $ anaemia
                                                                         <int> 0, 0, 0, 1, 1, 1, 1, 1, 0, 1,
## $ creatinine_phosphokinase <int> 582, 7861, 146, 111, 160, 47, 246, 31...
##$ diabetes
                                                               <int> 0, 0, 0, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0...
##$ejection fraction
                                        <int> 20, 38, 20, 20, 20, 40, 15, 60, 65, 3...
                                        <int> 1, 0, 0, 0, 0, 1, 0, 0, 0, 1, 1, 1, 0...
##$high blood pressure
## $ platelets
                                        <dbl> 265000, 263358, 162000, 210000, 32700...
##$ serum creatinine
                                        <dbl> 1.90, 1.10, 1.30, 1.90, 2.70, 2.10, 1...
## $ serum_sodium
                                        <int> 130, 136, 129, 137, 116, 132, 137, 13...
## $ sex
                                        <int> 1, 1, 1, 1, 0, 1, 1, 1, 0, 1, 1, 1, 1...
## $ smoking
                                        <int> 0, 0, 1, 0, 0, 1, 0, 1, 0, 1, 1, 1, 1, 0...
## $ time
                                        <int> 4, 6, 7, 7, 8, 8, 10, 10, 10, 10, 10, ...
## $ DEATH EVENT
                                        <int> 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1...
```

Let us summarise the data and note observations

### summary of data

summary(	data)								
##	age	ana	emia	creatinine_phosphokir		nase			
## Min.	:40.00	Min.	:0.0000	Min.	: 23.0				
## 1st Qu.	:51.00	1st Qu.:	0.0000	1st Qu.: 116.5					
## Mediar	า :60.00	Median	:0.0000	Median: 250.0					
## Mean	:60.83	Mean	:0.4314	Mean	: 581.8				
## 3rd Qu	.:70.00	3rd Qu.:	1.0000	3rd Qu.: 582.0					
## Max.	:95.00	Max.	:1.0000	Max.	:7861.0				
##	diabetes	ejection	_fraction hi	igh_blood_pressure		plat	elets		
## Min.	:0.000	00Min.	:14.00	Min.	:0.0000	Min.	: 25100		
## 1st Qu.	:0.0000	1st Qu.:	30.00	1st Qu.:0.0000		1st Q	1st Qu.:212500		
## Mediar	n :0.0000	Median	:38.00	Median :0.0000		Media	n :262000		
## Mean	:0.4181	Mean	:38.08	Mean	:0.3512	Mean	:263358		
## 3rd Qu	.:1.0000	3rd C	u.:45.00	3rd Qu	ı.:1.0000	3rd Q	u.:303500		

## Max.	:1.0000	Max.	:80.00	Max.	:1.000	0 1	Max.	:850000
## serum_cre	reatinine serum_sodium		l	sex		smoking		
## Min.	:0.500	Min.	:113.0	Min.	:0.0000	Min.	:0.000	0
## 1st Qu.:0.9	00	1st Qu.	:134.0	1st Qu.:0	.0000	1st Qu.	:0.0000	
## Median :1.100		Median :137.0		Median :1.0000		Median :0.0000		
## Mean	:1.394	Mean	:136.6	Mean	:0.6488	Mean	:0.321	1
## 3rd Qu.:1.400 3rd Qu.:14		:140.0	3rd Qu.:1	1.0000	3rd Qu.:1.0000			
## Max.	:9.400	Max.	:148.0	Max.	:1.0000	Max.	:1.000	0
## tir	## time DEATH_EVENT		VENT					
## Min. : 4.0 Min. :0.0000 ## 1st Qu.: 73.0		u.: 73.0						
1st Qu.:0.0000 ## Median :115.0 Median			Median					
:0.0000 ## Mean :130.3 Mean :0.3211 ##			3211 ##					
3rd Qu.:203.	3rd Qu.:203.0 3rd Qu.:1.0000 ## Max.		# Max.					
:285.0 Max. :1.0000								

#### Observations -

- 1. 299 observations for 13 variables
- 2. Age is between 40 and 95 so not much outliers by intuition
- 3. Death\_event should be converted to factor variable as they take only 2 values
- 4. Creatinine phosphokinase, platelets clearly has an outlier from max value which we will confirm later byunivariate analysis

## Missing/ NAs check

data2 <- na.omit(data)

'data2' has same rows as 'data' so there are no missing values in data

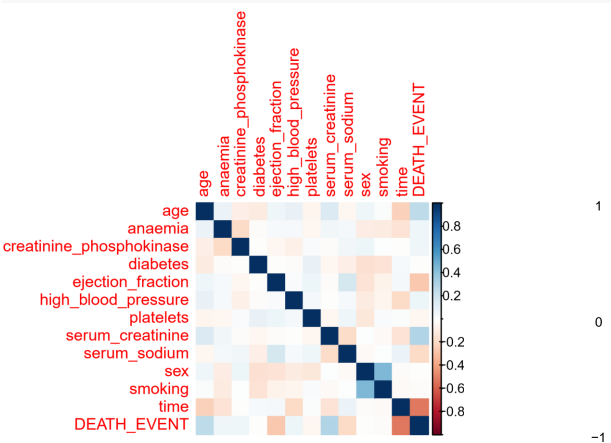
### **Correlation Plot**

M<-cor(data) head(round(M,2))

##					age ana	omia cros	atinina n	hosnhok	inase dial	hatas
## age	1.00	0.09	-0.08	-0 10 #	age ana # anaemi		1.00	-0.19	-0.01	octes
_							1.00	-0.19		
## creat	inine_ph	osphokir	nase -0.0	8	-0.1	9			1.00	-0.01
## diabe	etes	-0.10	-0.01	-0.01	1.00 ##	ejection	_fraction	0.06	0.03	-0.04
	0.00 ##	high_blo	od_press	sure	0.09	0.04	-0.07	-0.01#	# ejectior	_fraction
high_blo	ood_pres	sure pla	telets							
## age						0.06			0.09	-0.05
## anae	mia					0.03			0.04	-0.04
## creat	inine_ph	osphoki	nase			-0.04			-0.07	0.02
## diabe	etes					0.00			-0.01	0.09
## eject	ion_fract	tion				1.00			0.02	0.07
## high_	_blood_p	ressure				0.02			1.00	0.05
## age						0.16	-	-0.05 0.0	)7	0.02 -0.22

## anaemia	0.05	0.04 -0.09	-0.11 -0.14 ##
## creatinine_phosphokinase	-0.02	0.06 0.08	0.00 -0.01
	serum_creatinine serum	_sodium	sex smoking time
## diabetes	-0.05	-0.09 -0.16	-0.15 0.03
## ejection_fraction	-0.01	0.18 -0.15	-0.07 0.04
## high_blood_pressure	0.00	0.04 -0.10	-0.06 -0.20
##	DEATH_EVENT		
## age	0.25		
m ugc	0.23		
## anaemia	0.07		
## creatinine_phosphokinase	0.06		
uu dialaasa	0.00		
## diabetes	0.00		
## ejection_fraction	-0.27		
## high_blood_pressure	0.08		

### corrplot(M, method="color")



We see that age, anameia, creatinine\_phosphokinase, high\_blood\_pressure, serum\_creatinine have +ve correlation with death\_event

We see that ejection\_fraction, platelets, serum\_sodium, and time have -ve correlation with death\_event

But we will need deeper analysis to confirm these relationships.

### **Data Information**

```
# Converting to factor (dependent variable) data$DEATH_EVENT <-
factor(data$DEATH_EVENT)
```

### #Let's check how many zeros are in dataset colSums(data==0)

emia creatinine_phosphokinase	ana	age	##
0	170	0	##
high_blood_pressure	ejection_fraction	diabetes	##
194	0	174	##
serum_sodium	serum_creatinine	platelets	##
0	0	0	##
time	smoking	sex	##
0	203	105	##
		DEATH_EVENT	##
		203	##

### # Let's check their proportion to dataset as well round(colSums(data==0)/nrow(data)\*100,2)

```
##
                                                       anaemia creatinine phosphokinase
                            age
                       0.00 ## diabetes ejection fraction high blood pressure
##
       0.00
                56.86
##
       58.19
                0.00
                        64.88 ## platelets serum_creatinine serum_sodium
##
       0.00
                0.00
                        0.00 ## sex
                                        smoking time
                                                                                      0.00
##
                          35.12
                                                       67.89
##
                  DEATH_EVENT
##
                          67.89
```

Smoking, High BP, Diabetes, Anaemia are over 50% while sex is below 35%

The Event rate of survival is ~67.9%

Let's classify independent variables into -

- 1. Categorical -> Anaemia, Diabetes, High\_blood\_pressure, Sex, Smoking
- 2. Numeric -> Age, Creatinine phosphokinase, Ejection fraction, Platelets, serum creatinine, serum sodium, time

We also see that

```
Sex - Gender of patient Male = 1, Female =0
Diabetes - 0 = No, 1 = Yes
Anaemia - 0 = No, 1 = Yes
High_blood_pressure - 0 = No, 1 = Yes
Smoking - 0 = No, 1 = Yes
DEATH EVENT - 0 = No, 1 = Yes
```

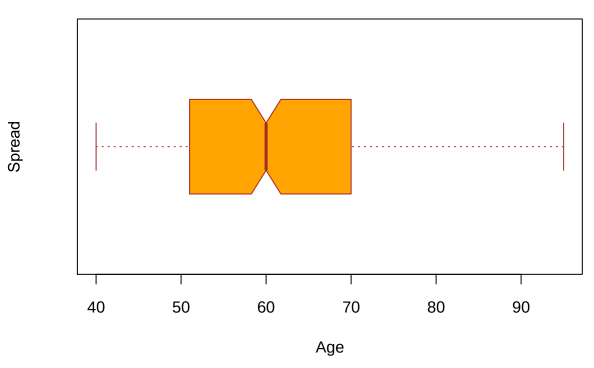
We note the scale of few variables like creatinine\_phosphokinase,platelets, ejection\_fraction and time. We can normalize the same before modeling but for now we will keep them as-is for the EDA.

### **Analysis**

Let's start with Outlier Analysis

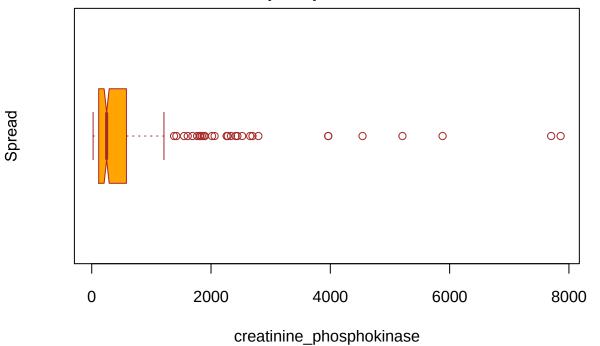
### Age

## **Age Box Plot**



We note no observed outlier in age creatinine\_phosphokinase

# creatinine\_phosphokinase Box Plot

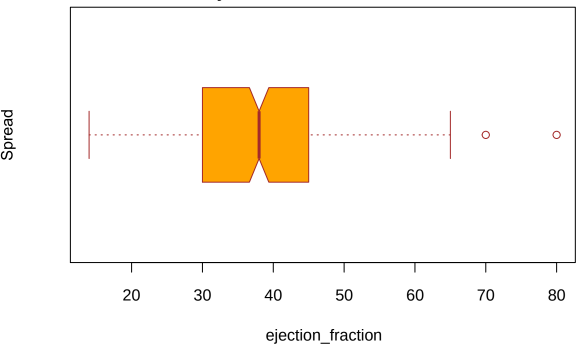


We notice some outliers at the positive side in creatinine\_phosphokinase with data above median more dispersed

### ejection\_fraction

```
boxplot(data$ejection_fraction, main =
    "ejection_fraction Box Plot", xlab =
    "ejection_fraction", ylab = "Spread", col =
    "orange", border = "brown", horizontal =
    TRUE,
    notch = TRUE
)
```

# ejection\_fraction Box Plot



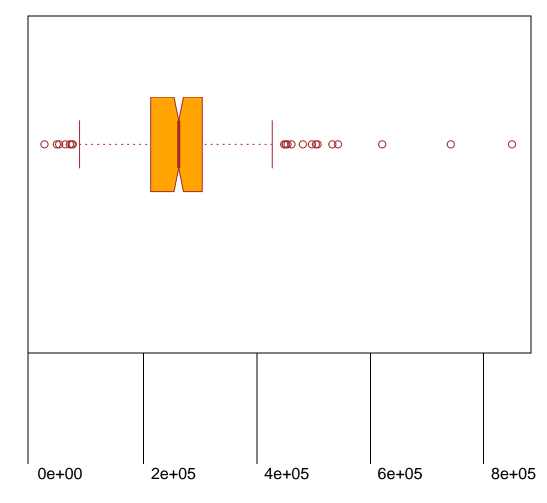
We notice 2 data points as outliers in ejection\_fraction

### platelets

```
boxplot(data$platelets, main = "platelets Box
    Plot", xlab = "platelets", ylab =
        "Spread", col = "orange", border =
        "brown", horizontal = TRUE,
        notch = TRUE
)
```

# platelets Box Plot





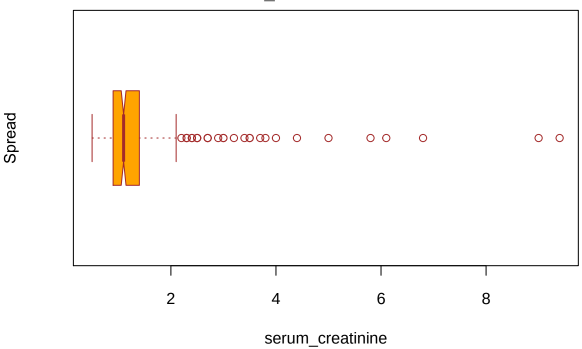
platelets

We notice outliers on both spectrum (high and low) in platelets

### serum\_creatinine

```
boxplot(data$serum_creatinine, main =
    "serum_creatinine Box Plot", xlab =
    "serum_creatinine", ylab = "Spread", col =
    "orange", border = "brown", horizontal =
    TRUE,
    notch = TRUE
)
```

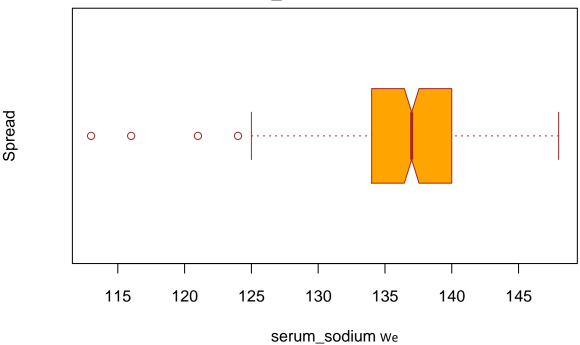
# serum\_creatinine Box Plot



We notice some outliers in serum\_creatinine on higher end (similar to creatinine\_phosphokinase). However these are in possible ranges medically.

### serum\_sodium

# serum\_sodium Box Plot

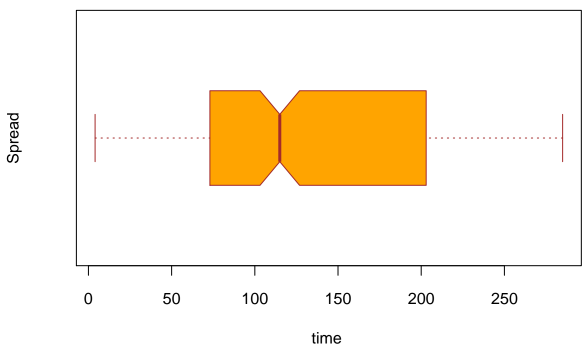


notice some outliers in serum\_sodium on lower end

### time

```
boxplot(data$time, main = "time Box
    Plot", xlab = "time ", ylab =
        "Spread", col = "orange",
        border = "brown", horizontal =
        TRUE,
        notch = TRUE
)
```





We notice no outliers in time (follow up period) however data above median is more dispersed

Note: While some of these are clear outliers, others must be checked with possible medical range

## Data Cleaning - Let's remove these outliers

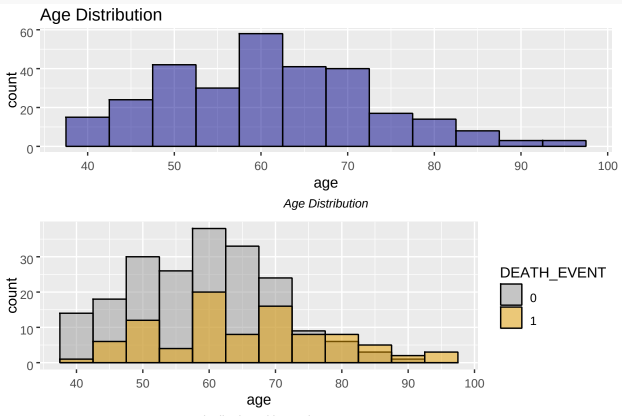
data <- data[data\$ejection\_fraction <70,] data <- data[data\$creatinine\_phosphokinase <7000,]

The new data now has only 295 observations - 4 observations were removed

## **Univariate analysis**

We will plot density distributions for numerical variables And check splits for categorical variables

#### Age



Age Distribution with Death Event

We see that as age increases, chances of death event go up as well We can also confirm this numerically;

```
# Let's create age ranges

data$age_tr[data$age < 50 & data$age >= 40]="40-50" data$age_tr[data$age < 60 & data$age >=

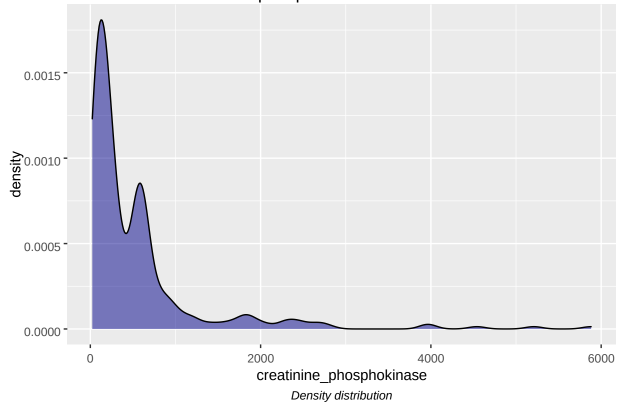
50]="50-60" data$age_tr[data$age < 70 & data$age >= 60]="60-70" data$age_tr[data$age < 80 &
```

```
data$age >= 70]="70-80" data$age_tr[data$age < 90 & data$age >= 80]="80-90"
data$age_tr[data$age < 100 & data$age >= 90]="90-100" table(data$DEATH_EVENT, data$age_tr)
##
##
            40-50 50-60 60-70 70-80 80-90 90-100
##
            35
                   62
                          66
                                 32
                                          6
                                                  1
      1
            10
                          27
                                 20
                                                  5
##
                   18
                                        13
```

Numerically, we can confirm the same observation (Higher death rate in higher ages)

### Creatinine\_phosphokinase

Distribution of creatinine phosphokinase



The distribution looks skewed

```
#let's create 10 splits of this variable
```

data\$creatinine\_phosphokinase\_tr <- cut(data\$creatinine\_phosphokinase, 10) table(data\$DEATH\_EVENT, data\$creatinine\_phosphokinase\_tr)

```
##
                (17.1,609] (609,1.19e+03] (1.19e+03,1.78e+03] (1.78e+03,2.37e+03]
##
## 0 156 24 7 7 ## 1 79 6 2 3
##
              (2.37e+03,2.95e+03] (2.95e+03,3.54e+03] (3.54e+03,4.12e+03]
##
##
      0
                              5
                                                                                1
                              1
      1
                                                       0
                                                                                1
##
##
##
              (4.12e+03,4.71e+03] (4.71e+03,5.3e+03] (5.3e+03,5.89e+03]
        0
                                 0 ##
##
                1
                         1
                                          1
                                                  0
                                                           0
                                                                   1
```

Numerically, we can see that for creatinine levels above 4000, death event seems to be higher but what about averages

```
aggregate(data[, c('creatinine_phosphokinase')], list(data$DEATH_EVENT), mean)

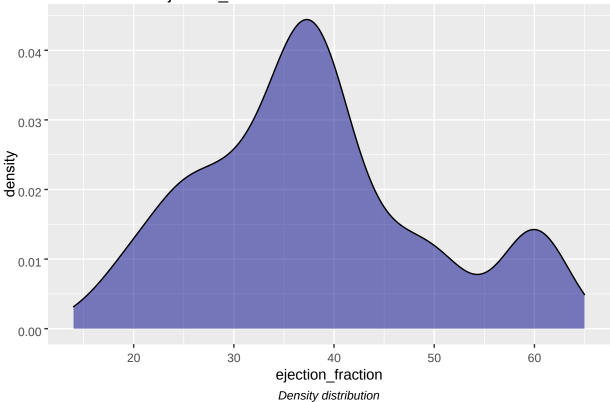
## Group.1 x ## 1 0

539.8465 ## 2 1 519.8817

Numerically, we can see that average creatinine levels are higher in case of death event
```

### Ejection\_fraction





The distribution has a major and a minor peak

```
#let's create 10 splits of this variable
```

data\$ejection\_fraction\_tr <- cut(data\$ejection\_fraction, 10) table(data\$DEATH\_EVENT, data\$ejection\_fraction\_tr)

```
##
##
                  (13.9,19.1] (19.1,24.2] (24.2,29.3] (29.3,34.4] (34.4,39.5]
        0
                 1
                          2
                                   18
                                            21
                                                     67 ##
                                                                       4
                                                                                16
##
                          21
                 13
##
                  (39.5,44.6] (44.6,49.7] (49.7,54.8] (54.8,59.9] (59.9,65.1]
##
##
      0
                    33
                                    15
                                                    15
                                                                                  28
##
      1
                      4
                                     5
                                                     6
                                                                    1
                                                                                    6
```

Numerically, we can see that ejection fraction is low in case of death event aggregate(data[, c('ejection\_fraction')],

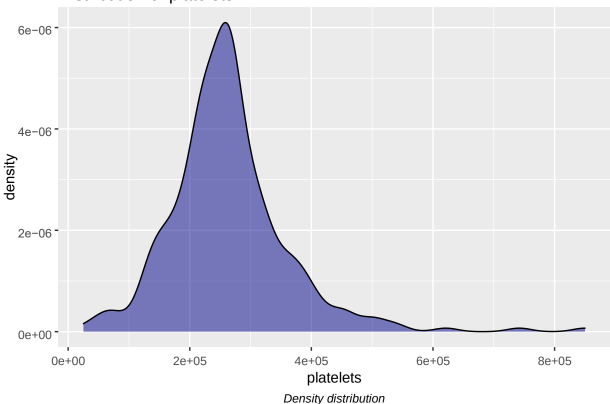
list(data\$DEATH\_EVENT), mean)

## Group.1 x ## 1 0 40.06931 ## 2 1 33.11828

Numerically, we can see that average ejection fraction is also low in case of death event platelets

ggplot(data,aes(x = platelets ))+geom\_density(fill = "dark blue", alpha = 0.5)+ labs(title = "Distribution of
platelets ", caption = "Density distribution")+ theme(plot.caption = element\_text(hjust = 0.5, face =
 "italic"))

### Distribution of platelets



The distribution looks pretty normal with some tail noticeable at the right extreme

#let's create 10 splits of this variable data\$platelets\_tr <cut(data\$platelets , 10) table(data\$DEATH\_EVENT, data\$platelets\_tr)

```
##
##
              (2.43e+04,1.08e+05] (1.08e+05,1.9e+05] (1.9e+05,2.73e+05]
      0
##
                                                                           91
                              5
      1
                                                    14
                                                                           43
##
##
              (2.73e+05,3.55e+05] (3.55e+05,4.38e+05] (4.38e+05,5.2e+05]
##
##
        0
                         18
                                 5 ##
                                          1
                                                  18
##
##
              (5.2e+05,6.03e+05] (6.03e+05,6.85e+05] (6.85e+05,7.68e+05]
      0
##
                                                                              1
                             0
                                                     1
                                                                              0
      1
##
##
##
          (7.68e+05,8.51e+05]
##
      0
                              1
##
      1
                              0
```

Numerically, we can see that platelets are low in case of death event

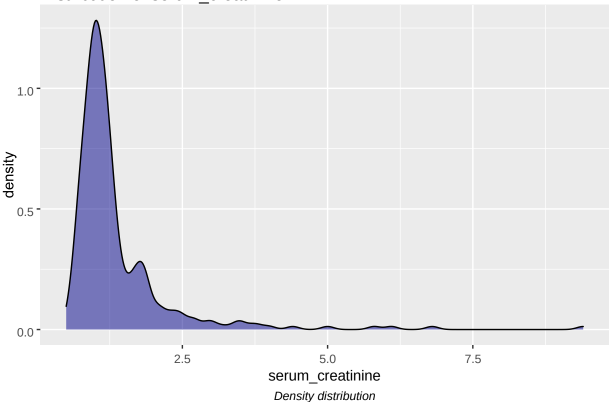
```
aggregate(data[, c('platelets')], list(data$DEATH_EVENT), mean)

## Group.1 x ## 1 0
266673.8 ## 2 1 256002.4
```

Numerically, we can see that average platelets are marginally lower in case of death event

### serum\_creatinine





The distribution looks similar to creatinine\_phosphokinase

```
#let's create 10 splits of this variable

data$serum_creatinine_tr <- cut(data$serum_creatinine , 10) table(data$DEATH_EVENT,
data$serum_creatinine_tr)
```

##

```
##
                   (0.491,1.39] (1.39,2.28] (2.28,3.17] (3.17,4.06] (4.06,4.95]
##
        0
                 168
                           23
                                   5
                                            4
                                                     0 ##
                                                              1
                                                                       47
                                                                                30
        9
                 3
                           1
##
                   (4.95,5.84] (5.84,6.73] (6.73,7.62] (7.62,8.51] (8.51,9.41]
##
##
                                   0
                                            0
                                                     0 ##
                                                                                0
         1
                 0
                           1
```

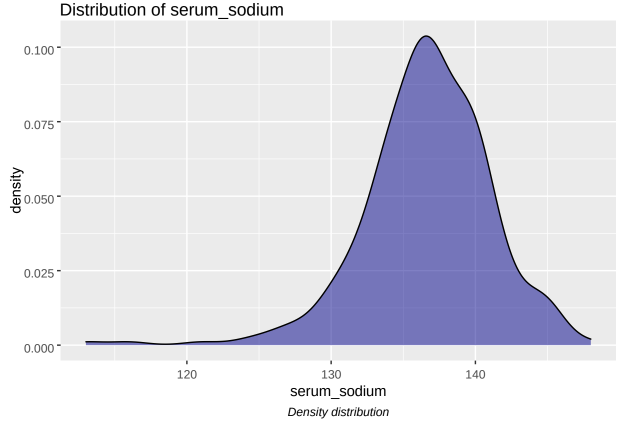
Numerically, we can see that death event is high when serum\_creatinine levels are above 1.39 and very high above 2.28

```
aggregate(data[, c('serum_creatinine')], list(data$DEATH_EVENT), mean)

## Group.1 x ## 1 0

1.184901 ## 2 1 1.775699
```

Numerically, we can see that average serum\_creatinine is high in case of death event **serum\_sodium** 



The distribution looks pretty normal with some tail noticeable at the left extreme

# #let's create 10 splits of this variable

```
data$serum_sodium_tr <- cut(data$serum_sodium , 10) table(data$DEATH_EVENT, data$serum_sodium_tr)
```

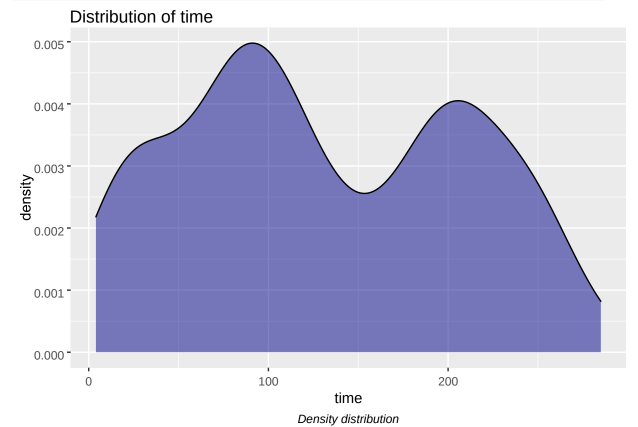
```
##
##
         (113,116] (116,120] (120,124] (124,127] (127,130] (130,134] (134,138]
##
      0
                               0
                                                                                31
                                                                                            69
                   1
                               0
                                            1
                                                        4
                                                                     6
                                                                                30
                                                                                            22
##
      1
##
##
         (138,141] (141,144] (144,148]
                                            7
##
      0
                  72
                              13
                                            4
      1
                  19
                               6
##
```

Numerically, we can see that serum\_sodium are low in case of death event

```
aggregate(data[, c('serum_sodium')], list(data$DEATH_EVENT), mean)
## Group.1 x ## 1  0
137.2178 ## 2  1 135.3118
```

Numerically, we can see that average serum sodium are marginally lower in case of death event time

```
ggplot(data,aes(x = time ))+geom_density(fill = "dark blue", alpha = 0.5)+ labs(title = "Distribution of time ",
    caption = "Density distribution")+ theme(plot.caption = element_text(hjust = 0.5, face = "italic"))
```



We note a bi-modal peak

```
#let's create 10 splits of this variable data$time tr <-
cut(data$time , 10) table(data$DEATH_EVENT, data$time_tr)
##
##
                   (3.72,32.1] (32.1,60.2] (60.2,88.3] (88.3,116] (116,144] (144,173]
## 0 5 4 33 32 14 13 ## 1 36 16 13 9 4 7
##
              (173,201] (201,229] (229,257] (257,285]
##
                                                         9
##
      0
                 28
                              33
                                           31
                               2
                                            2
                                                         0
      1
                   4
##
```

Numerically, we can see that follow up period was small in case of death event

```
aggregate(data[, c('time')], list(data$DEATH_EVENT), mean)

## Group.1 x

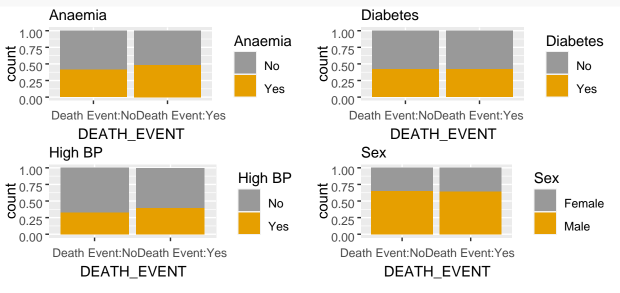
## 1 0 158.81188

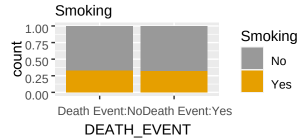
## 2 1 70.35484
```

Numerically, we can see that average follow up period is low in case of death event. This simply may illustrate that once deemed healthy, the patients may have stopped following up whereas diseased patients would undergo more checkups

#### Let's do EDA for categorical variables now

```
# Anaemia, Diabetes, High blood pressure, Sex, Smoking
a <- ggplot(data, aes(x = DEATH EVENT, fill = factor(anaemia)))+ geom bar(position = "fill")+
  scale_x_discrete(labels = c("Death Event:No","Death Event:Yes"))+ scale_fill_manual(values =
  c("#999999", "#E69F00"), name = "Anaemia", labels = c("No", "Yes"))+labs(subtitle = "Anaemia")
b<-ggplot(data, aes(x = DEATH_EVENT, fill = factor(diabetes)))+ geom_bar(position = "fill")+
  scale_x_discrete(labels = c("Death Event:No","Death Event:Yes"))+ scale_fill_manual(values = c("#999999",
  "#E69F00"), name = "Diabetes",
                                 labels = c("No","Yes"))+labs(subtitle = "Diabetes")
c<-ggplot(data, aes(x = DEATH_EVENT, fill = factor(high_blood_pressure)))+ geom_bar(position = "fill")+
  scale x discrete(labels = c("Death Event:No","Death Event:Yes"))+ scale fill manual(values = c("#999999",
  "#E69F00"), name = "High BP",
                                 labels = c("No", "Yes"))+labs(subtitle = "High BP")
d<-ggplot(data, aes(x = DEATH_EVENT, fill = factor(sex)))+ geom_bar(position = "fill")+
  scale x discrete(labels = c("Death Event:No","Death Event:Yes"))+
  scale_fill_manual(values = c("#999999", "#E69F00"), name = "Sex", labels =
  c("Female", "Male"))+labs(subtitle = "Sex")
e<-ggplot(data, aes(x = DEATH_EVENT, fill = factor(smoking)))+ geom_bar(position = "fill")+
  scale_x_discrete(labels = c("Death Event:No","Death Event:Yes"))+ scale_fill_manual(values =
  c("#999999", "#E69F00"), name = "Smoking", labels = c("No", "Yes"))+labs(subtitle = "Smoking")
grid.arrange(a,b,c,d,e)
```





We can see that Anaemia and High BP has significant difference for death event whereas others not so much

### **Tests**

### **Normality Test**

```
# univariate normality

data <- read.csv('/Users/mac/Downloads/heart_failure_clinical_records_dataset.csv') cm <-
colMeans(data) S <- cov(data)

d <- apply(data, MARGIN = 1, function(data)t(data - cm) %*% solve(S) %*% (data - cm))</pre>
```

Age

qqnorm(data[,"age"], main = "age") qqline(data[,"age"])

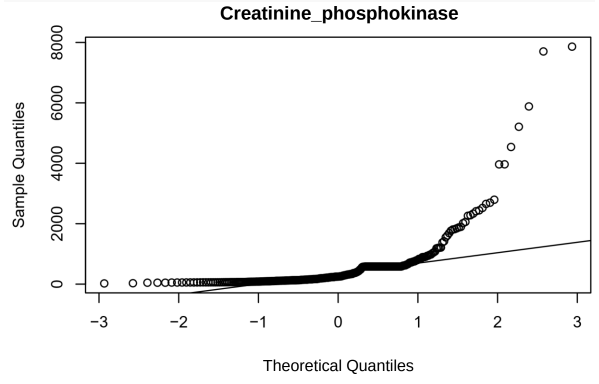


Age looks normally distributed

### Creatinine\_phosphokinase

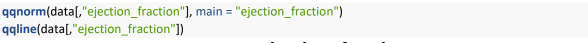
qqnorm(data[,"creatinine\_phosphokinase"], main = "Creatinine\_phosphokinase")

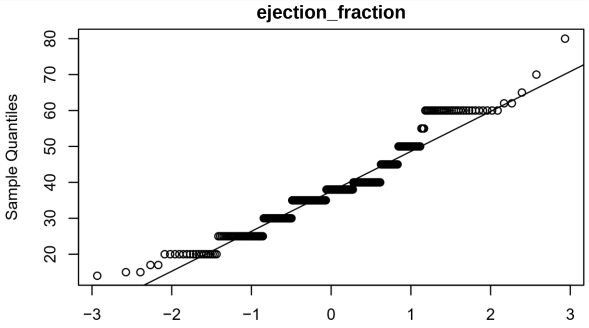




Creatinine\_phosphokinase doesn't looks normal but skewed

### ejection\_fraction



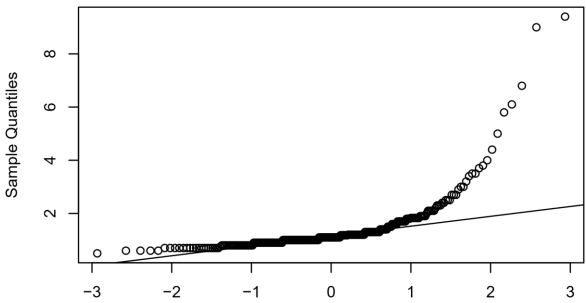


### **Theoretical Quantiles**

ejection\_fraction doeesn't look normal as well serum\_creatinine

```
qqnorm(data[,"serum_creatinine"], main = "serum_creatinine")
qqline(data[,"serum_creatinine"])
```

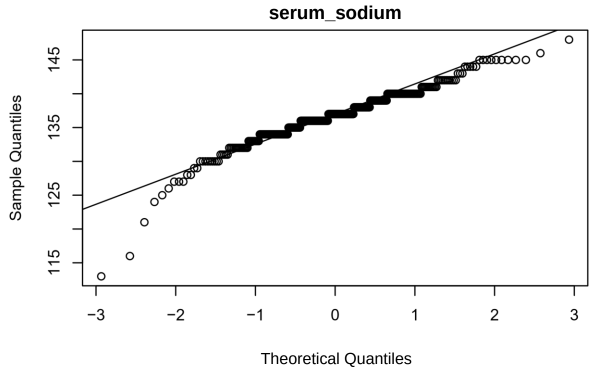
# serum\_creatinine



**Theoretical Quantiles** 

serum\_creatinine doesn't look normal but skewed serum\_sodium

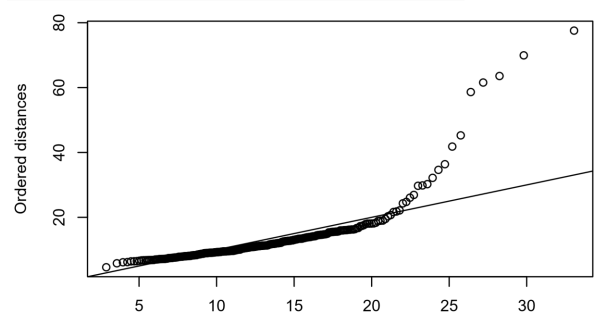
```
qqnorm(data[,"serum_sodium"], main = "serum_sodium")
qqline(data[,"serum_sodium"])
```



serum\_sodium looks normal but is slightly skewed on lower end

### Multi-variate normality

```
plot(qchisq((1:nrow(data) - 1/2) / nrow(data), df = 13), sort(d), xlab =
expression(paste(chi[13]^2, " Quantile")), ylab = "Ordered distances") abline(a = 0, b = 1)
```



## χ<sup>2</sup>13 Quantile

While plotting for multivariate normality, we see that data is non normal and has some skewness towards positive side

#### t-tests for death events vs not for each variable

```
# age
with(data,t.test(age[DEATH EVENT=="1"],age[DEATH EVENT=="0"],var.equal=TRUE))
## Two Sample t-test
## data: age[DEATH EVENT == "1"] and age[DEATH EVENT == "0"]
## t = 4.5206, df = 297, p-value = 8.917e-06
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## 3.643992 9.262758 ## sample estimates:
## mean of x mean of y
## 65.21528 58.76191
# anaemia
with(data,t.test(anaemia[DEATH_EVENT=="1"],anaemia[DEATH_EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test
##
## data: anaemia[DEATH_EVENT == "1"] and anaemia[DEATH_EVENT == "0"]
## t = 1.1446, df = 297, p-value = 0.2533
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -0.05057162 0.19117096 ## sample estimates:
## mean of x mean of y
## 0.4791667 0.4088670
# creatinine phosphokinase
with(data,t.test(creatinine_phosphokinase[DEATH_EVENT=="1"],
                    creatinine_phosphokinase[DEATH_EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test ##
## data: creatinine_phosphokinase[DEATH_EVENT == "1"] and creatinine_phosphokinase[DEATH_EVENT == "0"]
## t = 1.0832, df = 297, p-value = 0.2796
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -106.3109 366.5984 ## sample estimates:
## mean of x mean of y
## 670.1979 540.0542
# diabetes
with(data,t.test(diabetes[DEATH_EVENT=="1"], diabetes[DEATH_EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test
## data: diabetes[DEATH_EVENT == "1"] and diabetes[DEATH_EVENT == "0"]
```

```
## t = -0.033483, df = 297, p-value = 0.9733
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -0.1226917 0.1185866 ## sample estimates:
## mean of x mean of v
## 0.4166667 0.4187192
# ejection fraction
with(data,t.test(ejection_fraction[DEATH_EVENT=="1"],
                    ejection_fraction[DEATH_EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test ##
## data: ejection_fraction[DEATH_EVENT == "1"] and ejection_fraction[DEATH_EVENT == "0"]
## t = -4.8056, df = 297, p-value = 2.453e-06
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -9.580849 -4.013671 ## sample estimates:
## mean of x mean of y
## 33.46875 40.26601
#high blood pressure
with(data,t.test(high_blood_pressure[DEATH_EVENT=="1"],
                    high blood pressure[DEATH EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test
## data: high_blood_pressure[DEATH_EVENT == "1"] and high_blood_pressure[DEATH_EVENT == "0"]
## t = 1.3718, df = 297, p-value = 0.1711
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -0.03525453 0.19750823 ## sample estimates:
## mean of x mean of v
## 0.4062500 0.3251232
# platelets
with(data,t.test(platelets[DEATH_EVENT=="1"], platelets[DEATH_EVENT=="0"],var.equal=TRUE))
## Two Sample t-test
##
## data: platelets[DEATH EVENT == "1"] and platelets[DEATH EVENT == "0"]
## t = -0.84787, df = 297, p-value = 0.3972
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -34129.06 13576.17 ## sample estimates:
## mean of x mean of v
## 256381.0 266657.5
# serum creatinine
with(data,t.test(serum_creatinine[DEATH_EVENT=="1"],
                    serum creatinine[DEATH EVENT=="0"],var.equal=TRUE))
##
## Two Sample t-test
##
## data: serum_creatinine[DEATH_EVENT == "1"] and serum_creatinine[DEATH_EVENT == "0"]
```

```
## t = 5.3065, df = 297, p-value = 2.19e-07
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## 0.409539 0.892374 ## sample estimates:
## mean of x mean of v
## 1.835833 1.184877
# serum sodium
with(data,t.test(serum_sodium[DEATH_EVENT=="1"], serum_sodium[DEATH_EVENT=="0"],var.equal=TRUE))
## Two Sample t-test
##
## data: serum sodium[DEATH EVENT == "1"] and serum sodium[DEATH EVENT == "0"]
## t = -3.4301, df = 297, p-value = 0.0006889
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -2.8984440 -0.7850535 ## sample estimates:
## mean of x mean of y
## 135.3750 137.2167
# sex
with(data,t.test(sex[DEATH_EVENT=="1"], sex[DEATH_EVENT=="0"],var.equal=TRUE))
## Two Sample t-test
##
## data: sex[DEATH EVENT == "1"] and sex[DEATH EVENT == "0"]
## t = -0.074388, df = 297, p-value = 0.9408
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -0.1211614 0.1123355 ## sample estimates:
## mean of x mean of y
## 0.6458333 0.6502463
# smoking
with(data,t.test(smoking[DEATH_EVENT=="1"], smoking[DEATH_EVENT=="0"], var.equal=TRUE))
##
## Two Sample t-test
##
## data: smoking[DEATH_EVENT == "1"] and smoking[DEATH_EVENT == "0"]
## t = -0.21756, df = 297, p-value = 0.8279
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -0.1268080 0.1015617 ## sample estimates:
## mean of x mean of y
## 0.3125000 0.3251232
# time
with(data,t.test(time[DEATH_EVENT=="1"], time[DEATH_EVENT=="0"],var.equal=TRUE))
## Two Sample t-test
## data: time[DEATH EVENT == "1"] and time[DEATH EVENT == "0"]
## t = -10.686, df = 297, p-value < 2.2e-16
```

```
## alternative hypothesis: true difference in means is not equal to 0 ## 95 percent
confidence interval: ## -103.5612 -71.3478 ## sample estimates:
## mean of x mean of y ##
70.88542 158.33990
p-value is below 0.05 for 1.
age
2. serum_sodium
3. serum_creatinine
4. ejection_fraction5. time so we may conclude that death event does differ by these variables t-test multi-variate
# Hotelling's T2 test. Comparing multivariate means between death events and non-death event t2testdata <-
hotelling.test(age + anaemia + creatinine_phosphokinase + diabetes + ejection_fraction + high_blood_pressure+
platelets+serum_creatinine + serum_sodium + sex+smoking+time
                    ~ DEATH EVENT, data)
cat("T2 statistic =",t2testdatasstat[[1]],"\n")
## T2 statistic = 212.2908
print(t2testdata)
## Test stat: 17.036
## Numerator df: 12
## Denominator df: 286
## P-value: 0
The difference in means in the two groups taken together is significant as well
```

#### Homoskedasticity check

```
data$DEATH_EVENT <- factor(data$DEATH_EVENT) leveneTest(age ~
DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
             Df F value
                             Pr(>F)
## group
                17.1338 0.007981 **
##
           297
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
leveneTest(anaemia ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
              Df F value Pr(>F)
               1 1.3101 0.2533
## group
           297
##
leveneTest(creatinine_phosphokinase ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
              Df F value Pr(>F)
```

```
## group
                1 1.0303 0.3109
##
           297
leveneTest(diabetes ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
               Df F value Pr(>F)
## group
                1 0.0011 0.9733
##
           297
leveneTest(ejection_fraction ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
               Df F value Pr(>F)
##
## group
                1 3.7021 0.0553.
##
           297
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
leveneTest(high_blood_pressure ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
               Df F value Pr(>F)
## group
                1 1.8819 0.1711
           297
##
leveneTest(platelets ~ DEATH EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
        Df F value Pr(>F) ## group 1
        1.085 0.2984
           297
##
leveneTest(serum_creatinine ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
              Df F value
                               Pr(>F)
                1 16.242 7.087e-05 ***
## group
##
           297
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
leveneTest(serum_sodium ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
               Df F value Pr(>F)
                    5.274 0.02234 *
## group
             1
##
           297
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
leveneTest(sex ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
##
               Df F value Pr(>F)
               1 0.0055 0.9408
## group
```

```
297
##
```

```
leveneTest(smoking ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
               Df F value Pr(>F)
##
               1 0.0473 0.8279
## group
           297
##
leveneTest(time ~ DEATH_EVENT, data=data)
## Levene's Test for Homogeneity of Variance (center = median)
              Df F value
##
                             Pr(>F)
                1 7.9512 0.005129 **
## group
##
           297
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
p-value is below 0.05 for 1.
age
2. serum_creatinine
3. ejection fraction
4. time
5. serum_sodium
so we may conclude that variance between the two groups differ in them
One-way ANOVA tests: comparing univariate means
aov_age <- aov(age ~ DEATH_EVENT, data) summary(aov_age)
                      Df Sum Sq Mean Sq F value
                                                       Pr(>F)
                                              20.44 8.92e-06 ***
## DEATH EVENT
                           2714 2714.4
## Residuals
                    297 39449
                                   132.8
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
aov_anaemia <- aov(anaemia ~ DEATH_EVENT, data) summary(aov_anaemia)
##
        Df Sum Sq Mean Sq F value Pr(>F) ## DEATH EVENT 1
        0.32 0.3221
                        1.31 0.253
## Residuals
                    297 73.02 0.2459
aov_creatinine_phosphokinase <- aov(creatinine_phosphokinase ~ DEATH_EVENT, data)</pre>
```

1.173

```
summary(aov creatinine phosphokinase)
                  Df
                            Sum Sq Mean Sq F value Pr(>F)
```

1 ## Residuals 297 279450722 940912

## DEATH EVENT

aov\_diabetes <- aov(diabetes ~ DEATH\_EVENT, data) summary(aov\_diabetes)

1103933 1103933

0.28

```
##
                       Df Sum Sq Mean Sq F value Pr(>F)
## DEATH EVENT
                          0.00 0.00027
                                             0.001 0.973
## Residuals
                    297 72.74 0.24492
aov_ejection_fraction <- aov(ejection_fraction ~ DEATH_EVENT, data) summary(aov_ejection_fraction)
                     Df Sum Sq Mean Sq F value
##
                                                      Pr(>F)
## DEATH EVENT
                          3011 3011.4
                                             23.09 2.45e-06 ***
                    1
## Residuals
                                  130.4
                   297 38728
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' '1 aov_high_blood_pressure <- aov(high_blood_pressure ~ DEATH_EVENT,
data) summary(aov_high_blood_pressure)
                       Df Sum Sq Mean Sq F value Pr(>F)
##
## DEATH_EVENT
                           0.43 0.4290
                                             1.882 0.171
                    1
                    297 67.70 0.2279
## Residuals
aov_platelets <- aov(platelets ~ DEATH_EVENT, data) summary(aov_platelets)
##
                   Df
                          Sum Sq
                                        Mean Sq F value Pr(>F)
## DEATH_EVENT
                      1 6.883e+09 6.883e+09
                                                   0.719 0.397
## Residuals
                    297 2.844e+12 9.575e+09
aov_serum_creatinine <- aov(serum_creatinine ~ DEATH_EVENT, data) summary(aov_serum_creatinine)
##
                     Df Sum Sq Mean Sq F value
                                                      Pr(>F)
                                             28.16 2.19e-07 ***
## DEATH EVENT
                       1 27.62 27.618
## Residuals
                  297 291.30
                                  0.981
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
aov_serum_sodium <- aov(serum_sodium ~ DEATH_EVENT, data) summary(aov_serum_sodium)
##
                     Df Sum Sq Mean Sq F value
                                                      Pr(>F)
## DEATH EVENT
                    1
                            221 221.08
                                             11.77 0.000689 ***
## Residuals
                  297
                         5581
                                  18.79
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
aov sex <- aov(sex ~ DEATH EVENT, data) summary(aov sex)
                       Df Sum Sq Mean Sq F value Pr(>F)
## DEATH EVENT 1 0.00 0.00127 0.006 0.941 ## Residuals 297
68.13 0.22938
aov smoking <- aov(smoking ~ DEATH EVENT, data) summary(aov smoking)
                       Df Sum Sq Mean Sq F value Pr(>F)
##
## DEATH EVENT 1 0.01 0.01039 0.047 0.828 ## Residuals 297
65.17 0.21942
aov time <- aov(time ~ DEATH EVENT, data) summary(aov time)
```

```
##
                        Df Sum Sq Mean Sq F value Pr(>F)
## DEATH EVENT
                       1 498494 498494
                                               114.2 <2e-16 ***
## Residuals
                   297 1296647
                                     4366
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
p-value is below 0.05 for 1.
age
2. serum_creatinine
3. ejection fraction
4. time
5. serum sodium
so we may conclude that means between the two groups differ in them
```

#### Comparing multivariate means (One-way MANOVA)

```
mnv <- manova(as.matrix(data[,-13])~ DEATH_EVENT, data) summary(mnv)

## Df Pillai approx F num Df den Df Pr(>F)

## DEATH_EVENT 1 0.41684 17.036 12 286 < 2.2e-16 ***

## Residuals 297

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

We observe from MANOVA that estimated effects may be unbalanced indicating that mean between groups may be different

#### Multi-collinearity check

```
# Let us also look at Multicollinearity check
# Earlier we saw the correlation plot # We will check VIF for this purpose
# In classification, although linear regression isn't to be used
# For VIF, a rudimentary model lets us know the association
# between continuous and categorical variables data <-
read.csv('/Users/mac/Downloads/heart_failure_clinical_records_dataset.csv') mod <- Im( DEATH_EVENT ~
age+anaemia+creatinine_phosphokinase+ diabetes+ejection_fraction_high_blood_pressure+platelets+
serum_creatinine+serum_sodium+sex+smoking+time, data)
summary(mod)
##
## Call:
## Im(formula = DEATH EVENT ~ age + anaemia + creatinine phosphokinase +
##
                 diabetes + ejection_fraction + high_blood_pressure + platelets +
##
                serum_creatinine + serum_sodium + sex + smoking + time, data = data)
##
## Residuals:
```

```
##
                 1Q
                         Median 3Q
                                          Max ## -0.80866 -
        Min
0.28041 -0.04205 0.24742 0.96983
## Coefficients:
                                           Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                      1.664e+00 6.954e-01
                                                                   2.392 0.01738 *
## age
                                      5.767e-03 1.867e-03
                                                                   3.088 0.00221 **
## anaemia
                                        -2.766e-03 4.438e-02 -0.062 0.95035
## creatinine_phosphokinase 3.427e-05 2.247e-05
                                                                  1.525 0.12840
## diabetes
                                      1.928e-02 4.410e-02
                                                                  0.437 0.66236
                                        -9.834e-03 1.844e-03 -5.333 1.96e-07 ***
## ejection fraction
## high_blood_pressure
                                  -1.430e-02 4.565e-02 -0.313 0.75438
## platelets
                                  -8.370e-08 2.208e-07 -0.379 0.70492
## serum_creatinine
                                  8.527e-02 2.123e-02
                                                                    4.017 7.54e-05 ***
## serum_sodium
                                  -7.599e-03 5.024e-03 -1.513 0.13149
## sex
                                  -6.369e-02 5.108e-02 -1.247 0.21353
## smoking
                                  -5.733e-03 5.119e-02 -0.112 0.91091
                                  -2.733e-03 2.903e-04 -9.415 < 2e-16 ***
## time
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.3646 on 286 degrees of freedom
## Multiple R-squared: 0.4168, Adjusted R-squared: 0.3924 ## F-statistic: 17.04
on 12 and 286 DF, p-value: < 2.2e-16
vif(mod)
##
                                                          anaemia creatinine_phosphokinase
                              age
##
                        1.106067
                                                       1.087163
                                                                                      1.066014
##
                        diabetes
                                              ejection fraction
                                                                         high_blood_pressure
##
                        1.064324
                                                       1.067758
                                                                                      1.068377
```

serum creatinine

1.081241

smoking

1.285049

serum sodium

1.101927

1.138009

time

We see that most VIF values are below 1.5

##

## ##

##

This incdicates absence of multi-collinearity in our data

platelets

1.045809

1.337716

sex

### This concludes our initial EDA for the data