Lab 2: Statistical inference & hypothesis testing

Practice session covering topics discussed in Lecture 2

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GOAL OF TODAY'S PRACTICE SESSION

Consolidate understanding of inferential statistic, through R coding examples conducted on real biostatistics research data.

Lecture 2: topics

- Purpose and foundations of inferential statistics
- Getting to know the "language" of hypothesis testing
- Hypothesis testing
 - review examples
- A closer look at testing assumptions
 - more examples dealing with assumptions' violation

RENVIRONMENT SET UP & DATA

Needed R Packages

- We will use functions from packages base, utils, and stats (pre-installed and pre-loaded)
- We will also use the packages below (specifying package::function for clarity).

```
library(fs)
 5 library(here)
 6 library(janitor)
7 library(dplyr)
8 library(forcats)
9 library(tidyr)
12 library(BSDA)
13 library(rstatix)
14 library(car)
15 library(multcomp)
18 library(ggplot2)
19 library(ggstatsplot) # 'ggplot2' Based Plots with Statistical Details
20 library(ggpubr)
21 library(patchwork)
22 library(viridis)
23 library(ggthemes)
```

DATASETS FOR TODAY

For the most part, we will refer to a real clinical dataset (for which a *Creative Commons license* was granted) discussed in two articles (also open access):

- Ahmad, T., Munir, A., Bhatti, S. H., Aftab, M., & Raza, M. A. (2017). *Survival analysis of heart failure patients: A case study*. PLOS ONE, 12(7), e0181001. https://doi.org/10.1371/journal.pone.0181001
- Chicco, D., & Jurman, G. (2020). *Machine learning can predict survival of patients with heart failure from serum creatinine and ejection fraction alone*. BMC Medical Informatics and Decision Making, 20(1), 16. https://doi.org/10.1186/s12911-020-1023-5

Importing from project folder (previously downloaded

You can access the dataset either:

- From the UC Irvine Machine Learning Repository Heart Failure Clinical Records
- From the workshop website: use function here to specify the complete path of the input data folder



Make sure to match your own folder structure!

INSPECTING THE "HEART FAILURE" DATASET

What are the variables and their levels of measurem

The data, with medical records of **299 heart failure patient**, were collected at the Faisalabad Institute of Cardiology and at the Allied Hospital in Faisalabad (Punjab, Pakistan), during April–December 2015. See Table 1.

Table 1

Feature	Explanation	Measurement	Range
Age	Age of the patient	Years	[40,, 95]
Anaemia	Decrease of red blood cells or hemoglobin	Boolean	0, 1
High blood pressure	If a patient has hypertension	Boolean	0, 1
Creatinine phosphokinase	Level of the CPK enzyme in the blood	mcg/L	[23,, 7861]
(CPK)			
Diabetes	If the patient has diabetes	Boolean	0, 1
Ejection fraction	Percentage of blood leaving	Percentage	[14,, 80]
	the heart at each contraction		
Sex	Woman or man	Binary	0, 1
Platelets	Platelets in the blood	kiloplatelets/mL	[25.01,, 850.00]
Serum creatinine	Level of creatinine in the blood	mg/dL	[0.50,, 9.40]
Serum sodium	Level of sodium in the blood	mEq/L	[114,, 148]
Smoking	If the patient smokes	Boolean	0, 1
Time	Follow-up period	Days	[4,,285]
(target) death event	If the patient died during the follow-up period	Boolean	0, 1

mcg/L: micrograms per liter. mL: microliter. mEq/L: milliequivalents per litre

Look into the dataset just loaded in the R environment

Recall some base R functions from Lab 1

```
2 colnames(heart failure)
[1] "age"
                                 "anaemia"
[3] "creatinine phosphokinase" "diabetes"
[5] "ejection fraction"
                                 "high blood pressure"
    "platelets"
                                 "serum creatinine"
                                 "sex"
 [9] "serum sodium"
[11] "smoking"
                                 "time"
[13] "DEATH EVENT"
          2 nrow(heart failure)
[1] 299
          2 dim(heart failure)
[1] 299 13
```

Inspect the dataframe structure (base

R)

```
1 # What does the dataframe look like?
2 str(heart_failure)
```

```
'data.frame':
               299 obs. of 13 variables:
                                 75 55 65 50 65 90 75 60 65 80 ...
$ age
                           : num
$ anaemia
                                 0 0 0 1 1 1 1 1 0 1 ...
$ creatinine phosphokinase: int
                                 582 7861 146 111 160 47 246 315 157 123 ...
$ diabetes
                                 0 0 0 0 1 0 0 1 0 0 ...
$ ejection fraction
                                 20 38 20 20 20 40 15 60 65 35 ...
                          : int
$ high blood pressure
                                 1 0 0 0 0 1 0 0 0 1 ...
                          : int
$ platelets
                                 265000 263358 162000 210000 327000 ...
                                 1.9 1.1 1.3 1.9 2.7 2.1 1.2 1.1 1.5 9.4 ...
$ serum creatinine
$ serum sodium
                                 130 136 129 137 116 132 137 131 138 133 ...
  sex
                                 1 1 1 1 0 1 1 1 0 1 ...
$ smoking
                          : int 0 0 1 0 0 1 0 1 0 1 ...
$ time
                          : int 4 6 7 7 8 8 10 10 10 10 ...
                          : int 1 1 1 1 1 1 1 1 1 1 ...
$ DEATH EVENT
```

Inspect the dataframe structure (ski

Remember the skimr function skim?

```
1 # some variables
2 heart_failure %>% skimr::skim( age, DEATH_EVENT )
3
4 # the whole dataframe
5 heart_failure %>% skimr::skim()
```

```
You try...

Run skimr::skim() on your own either on the whole dataset or on any specific variable
```

notice there are no (missing values) NAs in any of the variables

Recode some variables for later ease of analy

I may need some variables coded as factor (e.g. categorical variables for plotting), and, while I am at it, I can add clearer labels for the variables' levels. Here, we are:

using tidyverse packages dplyr and forcats

female

105

male

194

adding new (recoded) variables called "oldname_f"

Some more dummy variables recoded as fact

[Mostly for illustration: it's totally fine (if not preferable) to keep these as binary [0,1] variables]

• It's worth learning the useful function dplyr::across¹, which allows to iteratively transform several columns at once!

```
2 fct cols = c("anaemia", "diabetes", "high blood pressure", "smoking")
  heart failure <- heart failure %>%
    dplyr::mutate(
      dplyr::across(
        .cols = all of(fct cols),
        # Functions to apply to each col
        .fns = ~as.factor (.x),
        .names = "{.col} f")) %>%
    dplyr::mutate(
      dplyr::across(
        .cols = ends with(" f") & !matches(c( "DEATH EVENT f", "sex f" )) ,
        # Functions to apply to each col(different syntax)
        .fns = ~forcats::fct recode(.x, yes = "1", no = "0" )))
```

1. This is a bit more advanced, but it will save a lot of typing in some situations...

(Small digression on dplyr::acros

Notice how dplyr::across(.cols = ..., .fns = ..., .names = ...) has these arguments:

- 1. cols = to select the columns which we want to transform (i.e. fct_cols)
 - with help from tidyselect functions: all_of, ends_with, and matches
- 2. $fns = \sim function(x)$ to specify the function
 - where ~function(.x) uses the "anonymous function" syntax of the tidyverse
 - and x inside the function is a "stand in" for each of the columns selected
- 3. [optional] names = to name the new cols created using {• col} in place of each of the transformed columns

VISUAL DATA **EXPLORATION FOR** THE "HEART FAILURE"

CONTINUOUS VARIABLES

Why is visual exploration important?

- Gaining insight on the variables (range, outliers, missing data)
- Preliminary check of assumptions for parametric hypothesis testing:
 - normally distributed outcome variables?
 - homogeneity of variance across groups?

Let's explore the **Heart failure dataset** with some data visualization...

- Following the referenced articles (which were mostly interested in predict mortality based on patients' characteristics), we will take the categorical, binary variable DEATH_EVENT_f as our main criterion to split the sample (into *survived* and *dead* patients) to explore any significant difference between groups in terms of means of known quantitative features.
- We will look at both:
 - continuous variables in the dataset (with the Probability Density Function (PDF))
 - discrete variables in the dataset (with the Probability Mass Function (PMF))

Age

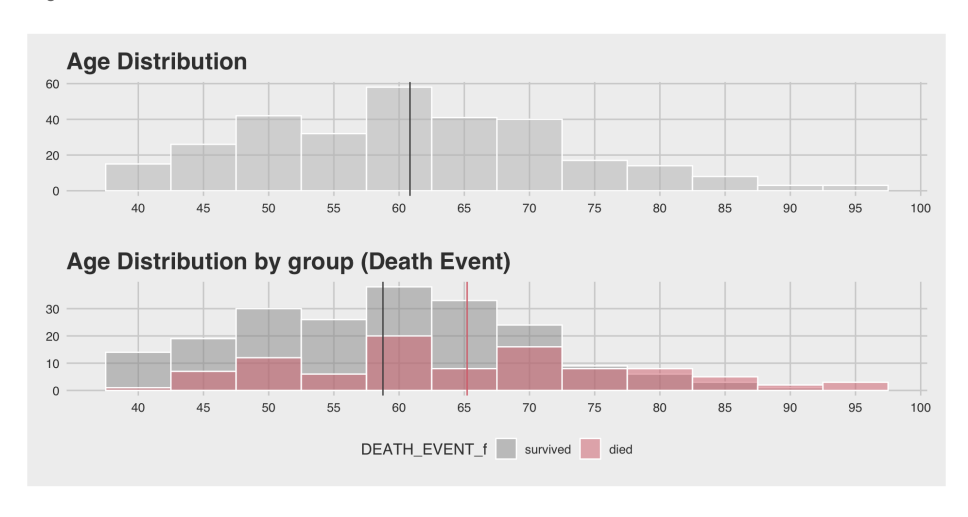
Introducing the handy R package patchwork which lets us compose different plots in a very simple and intuitive way

(check it out with ??patchwork)

```
1 age \leftarrow-ggplot(heart failure,aes(x = age ))+
     geom histogram(binwidth = 5, color = "white", fill = "grey",alpha = 0.5)+
     geom vline(aes(xintercept = mean(age)), color = "#4c4c4c")+
     theme fivethirtyeight()+
    labs(title = "Age Distribution" )+
     scale x continuous(breaks = seg(40,100,5))
   age2 <-ggplot(heart failure, aes(x = age, fill = DEATH EVENT f))+</pre>
     geom histogram(binwidth = 5, position = "identity", alpha = 0.5, color = "white")+
     geom vline(aes(xintercept = mean(age[DEATH EVENT == 0])), color = "#4c4c4c")+
     geom vline(aes(xintercept = mean(age[DEATH EVENT==1])), color = "#d8717b")+
     theme fivethirtyeight()+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     labs(title = "Age Distribution by group (Death Event)")+
     scale x continuous(breaks = seq(40,100,5))
18 library(patchwork) # The Composer of Plots
19 age + age2 + plot layout(ncol = 1)
```

Age

As the age increases, the incidence of death event seems to increase

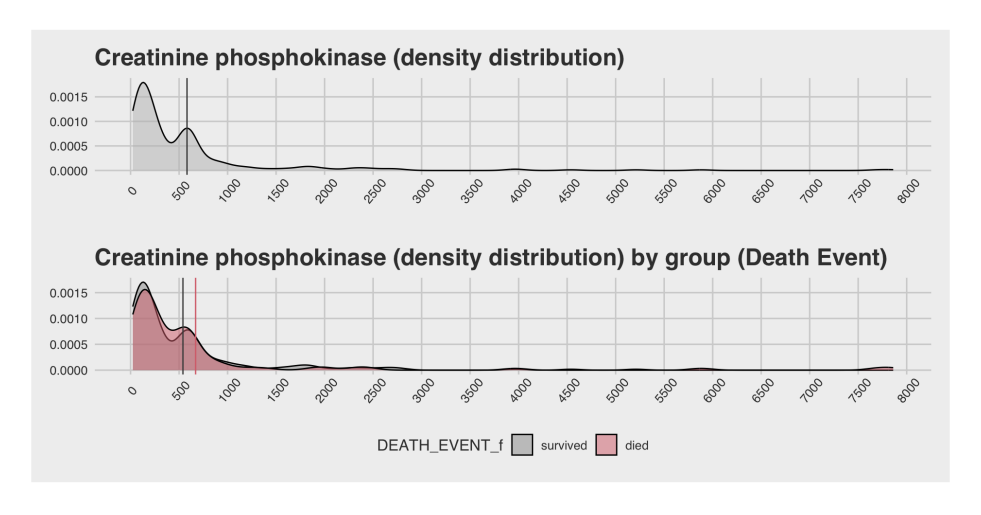


Creatinine Phosphokinase (CPK)

```
1 cpk <- ggplot(heart failure, aes(x = creatinine phosphokinase))+</pre>
     geom density(fill = "gray", alpha = 0.5)+
     scale x continuous(breaks = seq(0,8000, 500))+
     geom vline(aes(xintercept = mean(creatinine phosphokinase)), color = "#4c4c4c")+
     theme fivethirtyeight()+
     theme(axis.text.x = element text(angle=50, vjust=0.75))+
     labs(title = "Creatinine phosphokinase (density distribution)" )+
     theme(plot.caption = element text(hjust = 0.5, face = "italic"))
   cpk2 <- ggplot(heart failure, aes(x = creatinine phosphokinase, fill = DEATH EVENT f))+
     geom density(alpha = 0.5)+theme fivethirtyeight()+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     scale x continuous(breaks = seq(0,8000, 500))+
     geom vline(aes(xintercept = mean(creatinine phosphokinase[DEATH EVENT == 0])),
                color = "#4c4c4c") +
     geom vline(aes(xintercept = mean(creatinine phosphokinase[DEATH EVENT==1])),
                color = "#d8717b") +
     theme fivethirtyeight()+
     theme(axis.text.x = element text(angle=50, vjust=0.75))+
     labs(title = "Creatinine phosphokinase (density distribution) by group (Death Event)")
22 cpk + cpk2 + plot layout(ncol = 1)
```

Creatinine Phosphokinase (CPK)

This definitely doesn't look like a normal distribution!

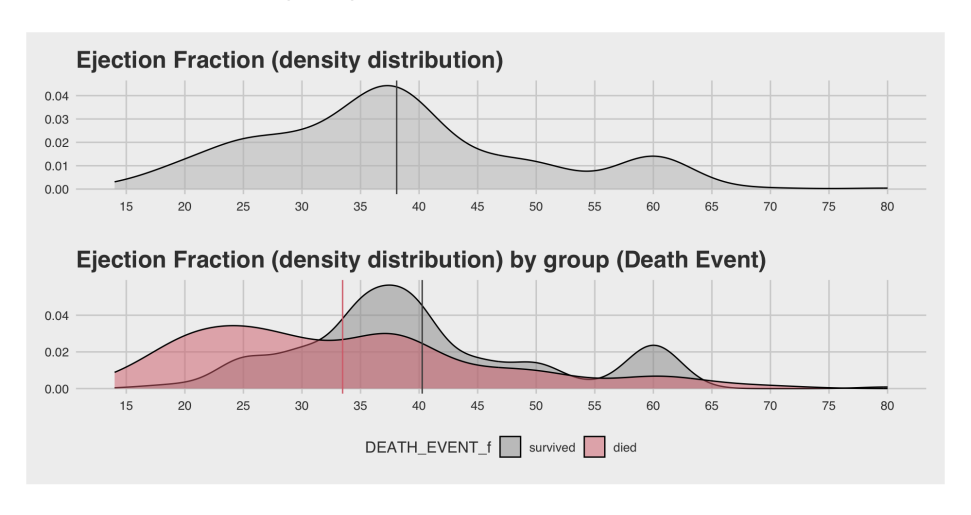


Ejection Fraction

```
1 ejf <- ggplot(heart failure,aes(x = ejection fraction))+</pre>
     geom density(fill = "gray", alpha = 0.5)+
     scale x continuous(breaks = seq(0,100, 5))+
     geom vline(aes(xintercept = mean(ejection fraction)), color = "#4c4c4c")+
     theme fivethirtyeight()+
     labs(title = "Ejection Fraction (density distribution)" )+
     theme(plot.caption = element text(hjust = 0.5, face = "italic"))
   ejf2 <- ggplot(heart failure, aes(x = ejection fraction, fill = DEATH EVENT f))+
     geom density(alpha = 0.5)+theme fivethirtyeight()+
     scale x continuous(breaks = seq(0,100, 5))+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     geom vline(aes(xintercept = mean(ejection fraction[DEATH EVENT == 0])),
                color = "#4c4c4c") +
     geom vline(aes(xintercept = mean(ejection fraction[DEATH EVENT==1])),
                color = "#d8717b") +
     labs(title = "Ejection Fraction (density distribution) by group (Death Event)")+
     theme fivethirtyeight()
20 ejf + ejf2 + plot layout(ncol = 1)
```

Ejection Fraction

This also doesn't look like a normal distribution... and there is a remarkable change in the *probability density function* (PDF) shape when we introduce the grouping variable

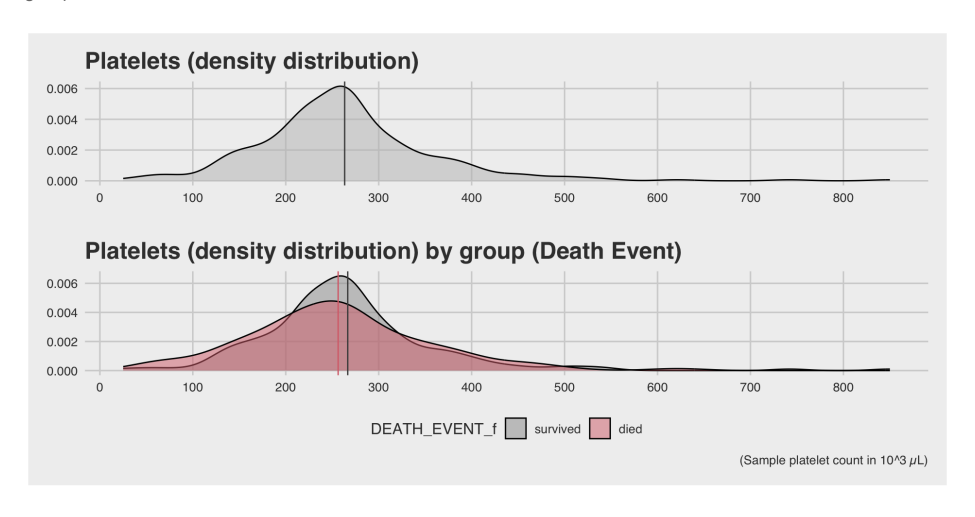


Platelets

```
2 heart failure <- heart failure %>% dplyr::mutate(plat norm = platelets/1000)
 4 plat <- ggplot(heart failure,aes(x = plat norm))+</pre>
     geom density(fill = "gray", alpha = 0.5)+
     scale x continuous(breaks = seq(0,800, 100))+
     geom vline(aes(xintercept = mean(plat norm)), color = "#4c4c4c")+
     theme fivethirtyeight()
     labs(title = "Platelets (density distribution)",
          y = "Density", x = "Sample platelet count (in 10^3 <math>\mu L)")
   plat2 <- ggplot(heart failure,aes(x = plat norm,fill = DEATH EVENT f))+</pre>
12
     geom density(alpha = 0.5)+theme fivethirtyeight()+
     scale x continuous(breaks = seq(0,800, 100))+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     geom vline(aes(xintercept = mean(plat norm[DEATH EVENT == 0])),
                 color = "#4c4c4c") +
     geom vline(aes(xintercept = mean(plat norm[DEATH EVENT==1])),
                 color = "#d8717b") +
     theme fivethirtyeight()
     labs(title = "Platelets (density distribution) by group (Death Event)",
21
22
          caption = "(Sample platelet count in 10^3 \mu L)")
24 plat + plat2 + plot layout(ncol = 1)
```

Platelets

Here the probability distributions resemble a Normal one and we observe more uniformity in the mean/variance across the 2 groups

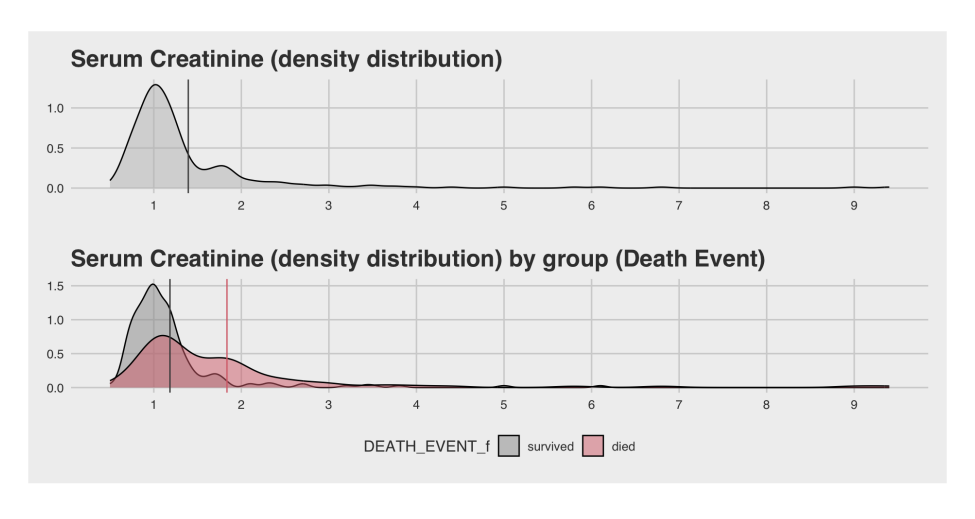


Serum Creatinine

```
1 ser cr <- ggplot(heart failure, aes(x = serum creatinine))+</pre>
     geom density(fill = "gray", alpha = 0.5)+
     scale x continuous(breaks = seq(0,10, 1))+
     geom vline(aes(xintercept = mean(serum creatinine)), color = "#4c4c4c")+
     theme fivethirtyeight()+
     labs(title = "Serum Creatinine (density distribution)" )+
     theme(plot.caption = element text(hjust = 0.5, face = "italic"))
   ser_cr2 <- ggplot(heart_failure,aes(x = serum creatinine,fill = DEATH EVENT f))+</pre>
     geom density(alpha = 0.5)+theme fivethirtyeight()+
     scale x continuous(breaks = seq(0,10,1))+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     geom vline(aes(xintercept = mean(serum creatinine[DEATH EVENT == 0])),
                color = "#4c4c4c") +
     geom vline(aes(xintercept = mean(serum creatinine[DEATH EVENT==1])),
                color = "#d8717b")+
     labs(title = "Serum Creatinine (density distribution) by group (Death Event)")+
     theme fivethirtyeight()
20 ser cr + ser cr2 + plot layout(ncol = 1)
```

Serum Creatinine

Another continuous random variable with a non-normal distribution (long right tails) and a seemingly important difference in variance between the groups.

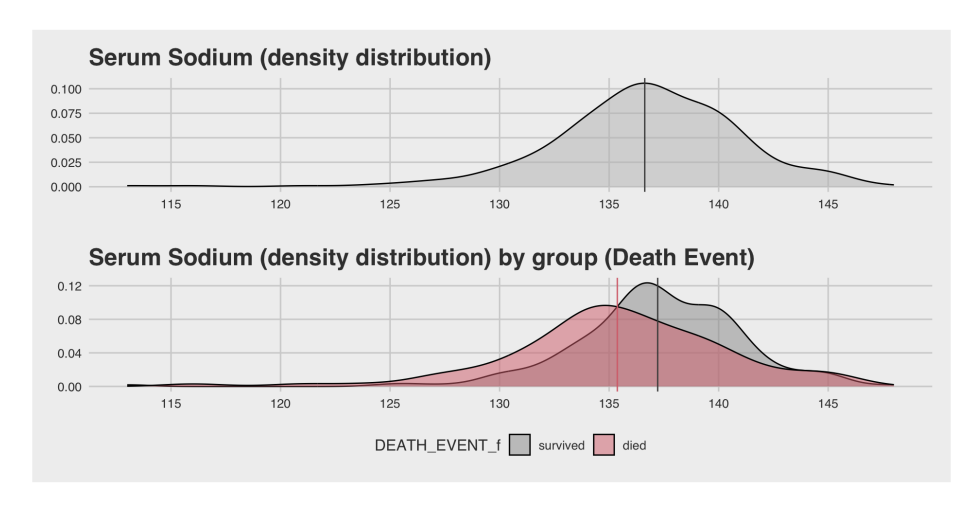


Serum Sodium

```
1 ser sod <- ggplot(heart failure,aes(x = serum sodium))+</pre>
     geom density(fill = "gray", alpha = 0.5)+
     scale x continuous(breaks = seq(0,150, 5))+
     geom vline(aes(xintercept = mean(serum sodium)), color = "#4c4c4c")+
    theme fivethirtyeight()+
    labs(title = "Serum Sodium (density distribution)" )
   ser sod2 <- ggplot(heart failure,aes(x = serum sodium,fill = DEATH EVENT f))+</pre>
     geom density(alpha = 0.5)+
     scale x continuous(breaks = seq(0,150, 5))+
     scale fill manual(values = c("#9999999", "#d8717b"))+
     geom vline(aes(xintercept = mean(serum sodium[DEATH EVENT == 0])),
                color = "#4c4c4c") +
     geom vline(aes(xintercept = mean(serum sodium[DEATH EVENT==1])),
                color = "#d8717b") +
     theme fivethirtyeight()+
     labs(title = "Serum Sodium (density distribution) by group (Death Event)")+
     theme fivethirtyeight()
20 ser sod + ser sod2 + plot layout(ncol = 1)
```

Serum Sodium

Same as above, except for the long left tails...



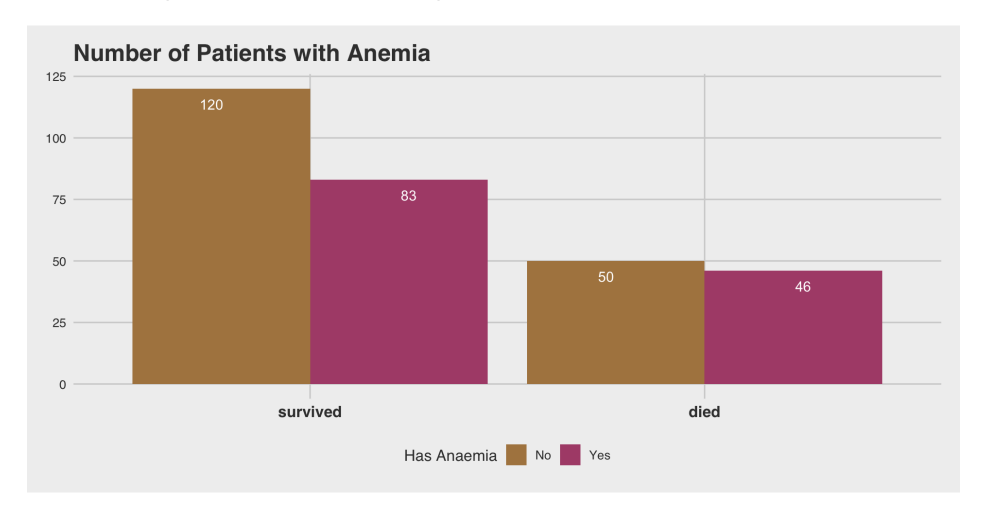
VISUAL DATA **EXPLORATION FOR** THE "HEART FAILURE"

DISCRETE VARIABLES

Anaemia

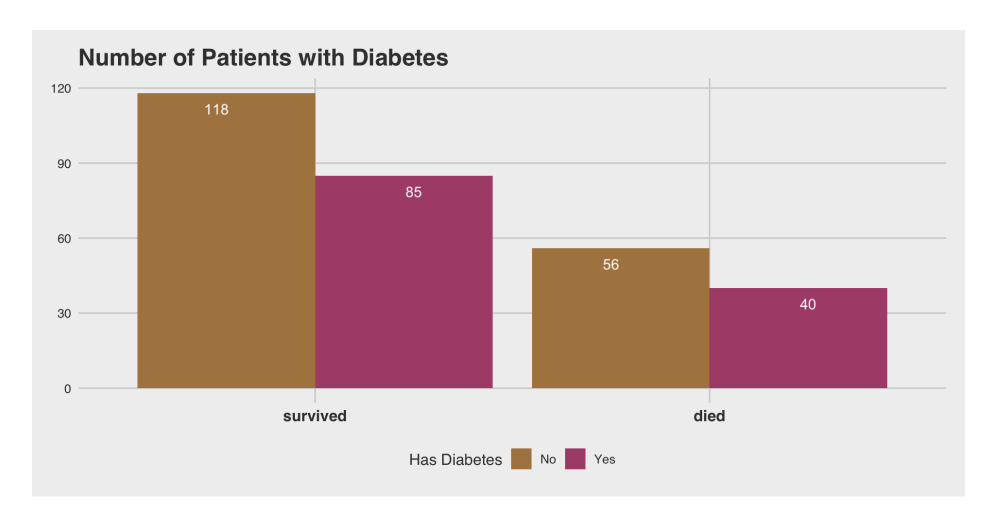
Anaemia

There seems to be a greater incidence of anaemia in group 'died'



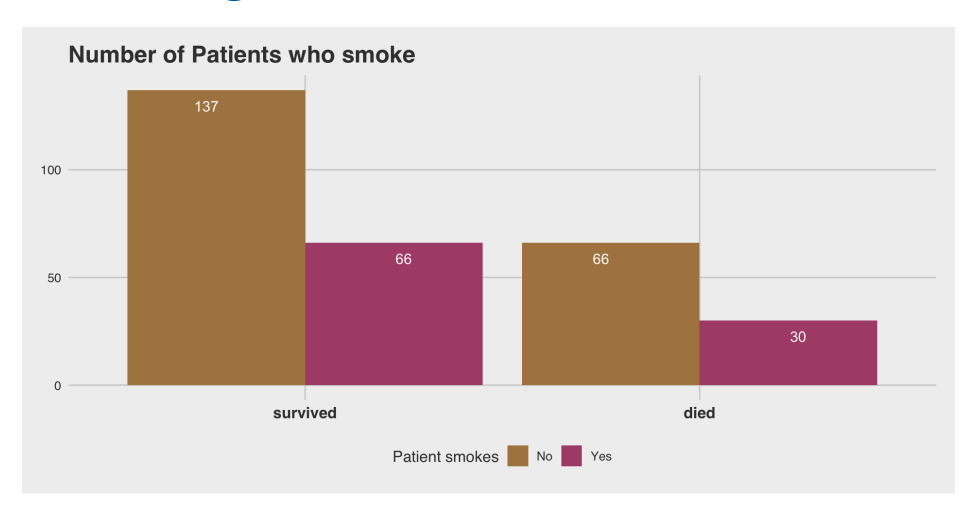
Diabetes

Diabetes



Smoking

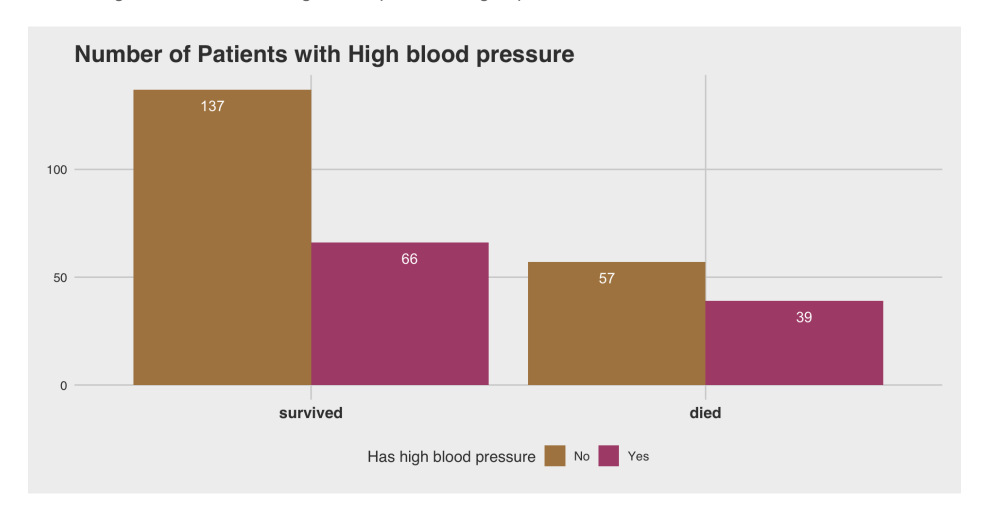
Smoking



High blood pressure

High blood pressure

There is also a greater incidence of high blood pressure in group 'died'



HYPOTHESIS TESTNG - some examples -

Let's continue to explore data from the **heart failure patients' dataset**, but this time using **hypothesis testing** as we learned in Lecture 2. We will do two types of test:

— EXAMPLE A —

(1 sample | n > 30 | Z test)

Comparing sample mean to a hypothesized population mean (with 2

Stating the above hypotheses more formally:

What is the population Total Platelet Count (TPC) mean for all people who suffered of heart failure (μ_{HF})?

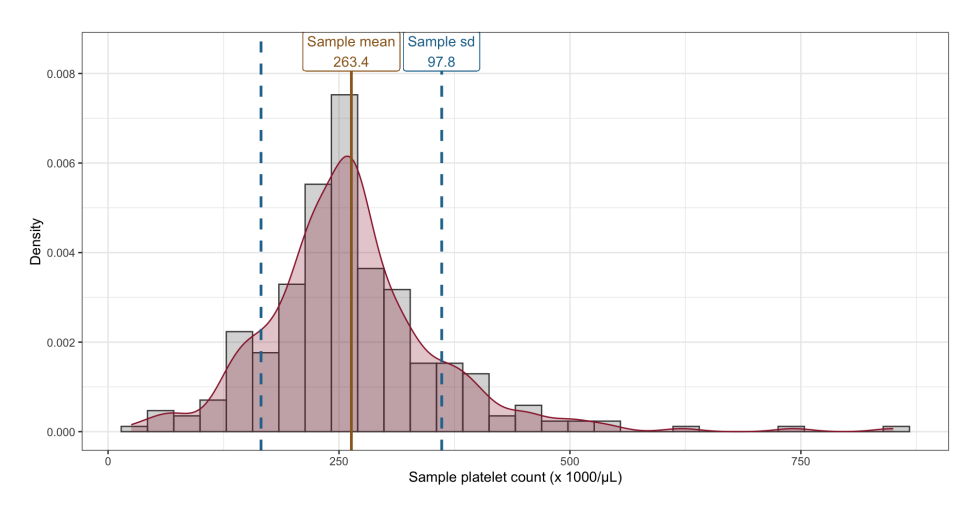
- ullet H_0 : there is no difference in mean TPC between patients who suffered heart failure and the general population
 - $\mu_{\rm HF}$ = 236 -> hypothesis of no effect or ("no difference")
- H_a : there is a difference in mean TPC between patients who have suffered heart failure and the general population ("some effect"). This can be formalized as either:
 - $\mu_{\rm HF}$ < 236 (one-sided test), or
 - $\mu_{\rm HF}$ > 236 (one-sided test), or
 - $\mu_{\rm HF} \neq 236$ (two-sided test)

1. Question: How does the mean platelets count in the patients' sample compare against a re

```
2 mean plat p <- round(mean(heart failure$plat norm), digits = 1)</pre>
3 sd plat p <- round(sd(heart failure$plat norm), digits = 1)</pre>
5 heart failure %>%
    ggplot(aes(x = plat norm))+
    geom histogram(aes(y = ..density..), bins=30, alpha=0.25, colour = "#4c4c4c") +
    geom density(colour = "#9b2339", alpha=0.25, fill = "#9b2339") +
    geom vline(xintercept = mean plat p, na.rm = FALSE, size = 1, color= "#9b6723") +
    geom vline(aes(xintercept = mean plat p + sd plat p),
                color = "#23749b", size = 1, linetype = "dashed") +
    geom vline(aes(xintercept = mean plat p - sd plat p),
                color = "#23749b", size = 1, linetype = "dashed") +
    geom label(aes(x=mean plat p, y=0.0085, label=paste0("Sample mean\n", mean plat p)),
                color = "#9b6723") +
    geom label(aes(x=361, y=0.0085, label=paste0("Sample sd\n", sd plat p)),
                color = "#23749b") +
    theme bw() + labs(y = "Density", x = "Sample platelet count (x 1000/\muL)")
```

1. Question: How does the mean platelets count in the patients' sample compare against a re

For a general population, the Total Platelet Count (TPL) has μ =236 (1000 / μ L) and σ = 59 (1000 / μ L). Below is the sample distribution:



2.a Computation of the test statistic

In this case, we have:

- a large sample (n > 100)
- a known σ^2 (of the reference population)
- the observed sample mean \bar{x} and sample sd s.

So we can compute:

$$Z_{\text{calc}} = \frac{\bar{x} - \mu}{\frac{\sigma}{\sqrt{n}}}$$

Let's do it "by hand" first to see the steps

```
1 # General Population of reference
2 mu <- 236
3 sigma <- 59
4 # Sample of HF patients
5 n <- 299
6 x_HF <- mean(heart_failure$plat_norm) # 263.358
7 s_HF <- sd(heart_failure$plat_norm) # 97.80424
8 # IF large sample & KNOWN pop variance
9 std_err_HF <- sigma /sqrt(n) # 3.412058
10 z_calc_HF <- (x_HF - mu) / std_err_HF # 8.018043</pre>
```

2.b Computation of the p-value associated to the test statistic

To find the **p-value** associated with a z-score in R, we can use the **pnorm()** function, which uses the following syntax:

- q: The z-score
- mean: The mean of the normal distribution. Default is 0.
- sd: The standard deviation of the normal distribution. Default is 1.
- lower tail:
 - If TRUE, the probability to the left of q in the normal distribution is returned
 - If FALSE, the probability to the right is returned. Default is TRUE.

```
1 # Left-tailed test
2 p_value_l <- stats::pnorm(z_calc_HF, mean = 0, sd = 1, lower.tail = TRUE)
3 # Right-tailed test
4 p_value_r <- stats::pnorm(z_calc_HF, mean = 0, sd = 1,lower.tail = FALSE)
5 # Two-tailed test (our case)
6 p_value_two <- 2*stats::pnorm(z_calc_HF, mean = 0, sd = 1, lower.tail = FALSE)</pre>
```

2.c Computation of the p-value associated to the test statistic

• <u>A</u> Let's see how this could be done using an R function BSDA::z.test

```
One-sample z-Test

data: heart_failure$plat_norm

z = 8.018, p-value = 1.074e-15

alternative hypothesis: true mean is not equal to 236

95 percent confidence interval:

256.6705 270.0455

sample estimates:

mean of x

263.358
```

Same results!

3. Results and interpretation

1. Based on the critical region, the calculated test statistic $z_calc_HF = 8.0180$ falls in the CRITICAL REGION (well beyond the critical point)

```
1  # given
2  z_critical <- c(-1.96, +1.96) # (Z score corresponding to α = 0.05)
3  # Check
4  z_calc_HF > z_critical
```

[1] TRUE TRUE

2. Based on the p-value, p_value_two = 1.07443e-15 is much much smaller than α

```
1 # Check
2 p_value_two < 0.05
```

[1] TRUE

DECISION: we reject the Null Hypothesis (basically we conclude that it is extremely unlikely that the sample we drew could have occurred just by chance). So the test indicates that, indeed, there is a difference between heart failure patients and the general population in terms of average platelets count.

— EXAMPLE B —

(1 sample | n < 30 | t test)

Comparing sample mean to a hypothesized population mean (with t test)

Same question, but with a *smaller sample* to work on (this varies, but generally it means n < 30). Imagine the patients were only observed over a **follow-up period of 21 days**, and also let's assume we don't know the population's variance

Stating the hypothesis more formally:

What is the population Total Platelet Count (TPC) mean for all people who suffered of heart failure (μ_{HF21d}) in the past 21 days or less?

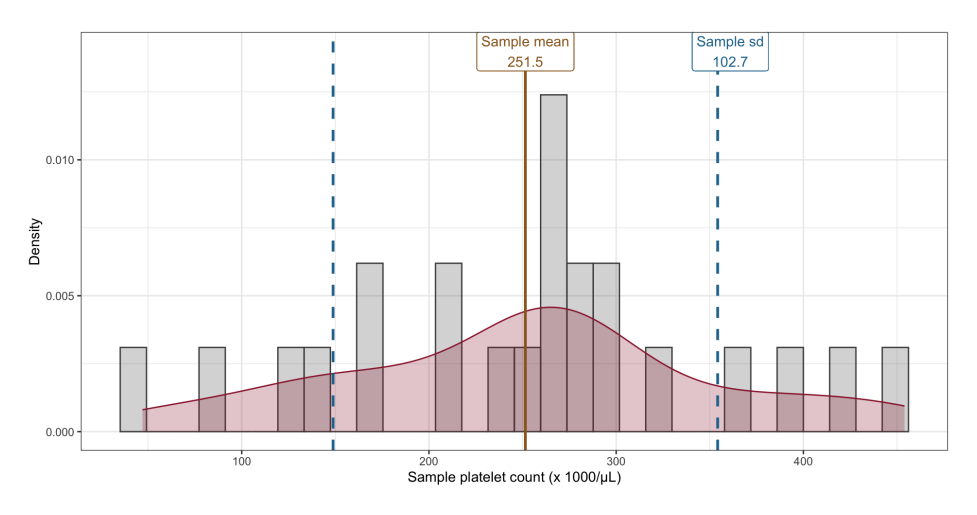
- ullet H_0 : there is no difference in mean TPC between patients who suffered heart failure (visited in 21 days) and the general population
 - $\mu_{\rm HF21d}$ = 236 -> hypothesis of no effect or ("no difference")
- H_a : there is a difference in mean TPC between patients who have suffered heart failure and the general population ("some effect"). This can be formalized as:
 - $\mu_{\rm HF21d} \neq 236$ (two-sided test)

1. Question: How does the mean platelets count in the patients' sample compare against a re

```
2 heart 21d <- heart failure %>% dplyr::mutate(plat norm = platelets/1000) %>%
3 filter(time <= 21)</pre>
5 mean plat p <- round(mean(heart 21d$plat norm), digits = 1)
6 sd plat p <- round(sd(heart 21d$plat norm), digits = 1)
8 heart 21d %>%
     ggplot(aes(x = plat norm))+
     geom histogram(aes(y = ..density..), bins=30, alpha=0.25, colour = "#4c4c4c") +
     geom density(colour = "#9b2339", alpha=0.25, fill = "#9b2339") +
     geom vline(xintercept = mean plat p, na.rm = FALSE, size = 1, color= "#9b6723") +
     geom vline(aes(xintercept = mean plat p + sd plat p),
                color = "#23749b", size = 1, linetype = "dashed") +
     geom vline(aes(xintercept = mean plat p - sd plat p),
                color = "#23749b", size = 1, linetype = "dashed") +
     geom label(aes(x=mean plat p, y=0.014, label=paste0("Sample mean\n", mean plat p)),
                color = "#9b6723") +
     geom label(aes(x=361, y=0.014, label=paste0("Sample sd\n",sd plat p)),
22
                color = "#23749b") +
     theme bw() + labs(y = "Density", x = "Sample platelet count (x 1000/\muL)")
```

1. Question: How does the mean platelets count in the patients' sample compare against a re

For a general population, the Total Platelet Count (TPL) has μ =236 (1000 / μ L) and σ = 59 (1000 / μ L). Below is the smaller sample distribution:



2.a Picking the suitable test

In this case, we have:

- a "small" sample n = 23
- an unknown σ^2 (of the reference population)
- We obtained the sample mean \bar{x} and sample sd s.

So we can compute:

$$t_{\text{calc}} = \frac{\bar{x} - \mu}{\frac{s_{\bar{x}}}{\sqrt{n-1}}}$$

2.b Computation of the test statistic

Option 1: Let's compute the t test "by hand"

Option 2: Let's compute the t test with stats::t.test

2.c Computation of the p-value associated to the test star

Option 1: "by hand"

To find the **p-value** associated with a t-score in R, we can use the pt(q, df, lower.tail = TRUE) function, which uses the following syntax:

- q: The t-score
- df: The degrees of freedom
- lower.tail:
 - TRUE to calculate the probability to the left of q which is called as left-tailed test
 - FALSE as right-tailed test.

```
1 # ---- Option 1
2 # -- Left-tailed test
3 #pt(t_stat_HF_21d, df_HF_21d, lower.tail = TRUE)
4
5 # -- Right-tailed test
6 #pt(t_stat_HF_21d, df_HF_21d, lower.tail = FALSE)
7
8 # -- Two-tailed test (our case)
9 p_value_t_test <- 2*pt(t_calc, df_HF_21d, lower.tail = FALSE) # 0.4863214</pre>
```

Option 2: from results of stats::t.test A

```
1 # ---- Option 2
2 # extract p_value from results df
3 p_value_v2 <- t_stat_HF_21d_v2[["p.value"]] # 0.4766892
```

3. Results and interpretation

1. Based on the critical region, $t_calc \approx 0.71$ is smaller than the t critical value, i.e. it falls within the region of acceptance, so he null hypothesis is not rejected

```
1 #find two-tailed t critical values
2
3 t_crit_two <- qt(p=.05/2, df=22, lower.tail=FALSE) # 2.073873
4 # Compare t score against t critical
5 t_calc > t_crit_two # FALSE
[1] FALSE
```

2. Based on the p-value, p_value \simeq 0.48 is larger than α , i.e. the probability of observing a test statistic (assuming H_0 is true) is quite large

```
1 # Check
2 p_value_t_test < 0.05 # FALSE
[1] FALSE</pre>
```

DECISION: we FAIL to reject H_0 . So the test indicates that there is not a statistically significant difference between heart failure patients visited within 21 days and the general population in terms of average platelets count.



Note

What changed testing a sample with smaller n, instead of a large one?

- EXAMPLE C -

(2 samples | t test)

Comparing two independent sample means (t t

This time, we investigate if there might be an actual difference in the Platelet Count means **between the patients who died and the patients who survived** heart failure.

Stating the above hypotheses more formally:

Is there a statistically significant difference between the mean values of two groups?

- ullet H_0 : The two population means are equal
 - $\mu_1 = \mu_0 \iff \mu_1 \mu_0 = 0$
- H_a : There is a mean difference between the two groups in the population. Possible directional difference formulation (two-tailed, left-tailed, right-tailed)
 - $\mu_1 \neq \mu_0 \iff \mu_1 \mu_0 \neq 0$ (the two population means are not equal)
 - $\mu_1 < \mu_0 \iff \mu_1 \mu_0 < 0$ (population 1 mean is less than population 0 mean)
 - $\mu_1 > \mu_0 \Longleftrightarrow \mu_1 \mu_0 > 0$ (population 1 mean is greater than population 0 mean)

Comparing two independent sample means (t test) (c

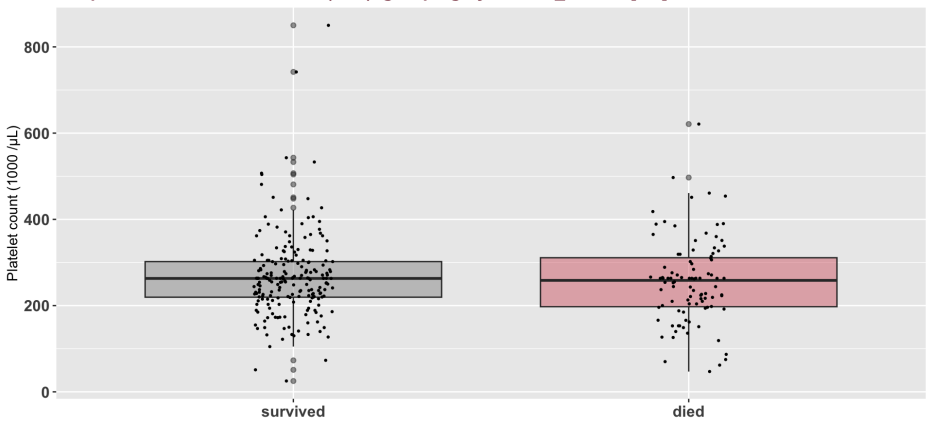
1. Question: Is there a statistically significant difference between the Platelet Counts in the patients

```
1  # boxplot by group
2  heart_failure %>%
3  ggplot(mapping = aes(y = plat_norm, x = DEATH_EVENT_f, fill = DEATH_EVENT_f)) +
4  geom_boxplot(alpha=0.5) +
5  #geom_violin(alpha=0.5) +
6  geom_point(position = position_jitter(width = 0.1), size = 0.5)+
7  scale_fill_manual(values = c("#999999", "#d8717b")) +
8  # drop legend and Y-axis title
9  theme(plot.title = element_text(size = 14,face="bold", color = "#873c4a"),
10  legend.position = "none",
11  axis.text.x = element_text(size=12,face="bold"),
12  axis.text.y = element_text(size=12,face="bold")) +
13  labs(title = "Boxplot of Total Platelet Count (TPL), grouping by DEATH_EVENT [0,1]",
14  x = "", y = "Platelet count (1000 /µL)")
```

Comparing two independent sample means (t test) (c

There seems to be no major difference in the two groups

Boxplot of Total Platelet Count (TPL), grouping by DEATH_EVENT [0,1]



2. Verify the assumptions for independent t-test

- 1. The 2 samples ("died" and "survived") must be independent V
- 2. The dependent variable is scaled in intervals (Platelets Count in 10³ "/ μ L")
- 3. The dependent variable is normally distributed (Platelets Count in 10³ "/ μ L")
- (If not, use *non parametric* test)
- 4. The variance within the 2 groups should be similar ?
- (If not, perform Welch's t-test)

Preliminary Fisher's F test to check for variance equality

We can compute the Fisher test "by hand"

[1] 1.020497

Preliminary Fisher's F test to check for variance equality (.cont)

```
1 ## -- Define the critical value of F distribution for a risk of alpha = 0.05
2 # qf(p=.05, df1 = n_died-1, df2 = n_survived-1, lower.tail = FALSE) # RIGHT-Tailed
3 # qf(0.95, df1 = n_died-1, df2 = n_survived-1, lower.tail = FALSE) # LEFT- Tailed
4 qf(c(0.025, 0.975), df1 = n_died-1, df2 = n_survived-1) # TWO-Tailed

[1] 0.6994659 1.3987233

1 ## --Compute the exact p-value (two-tailed)
2 p_value_f <- 2 * (1 - pf(F_ratio, df1 = (n_died-1), df2 = (n_survived-1)))
3 p_value_f</pre>
[1] 0.8914982
```

A test statistic (F) of 1.02 is obtained, with degrees of freedom 95 and 202.

The p-value is 0.89, greater than the p-value threshold of 0.05. This suggests we can not reject the null hypothesis of equal variances.

The variance within the 2 groups should be similar $\sqrt{\ }$ -> we can run a t-test.

3.a Computation of t test statistic

Since we verified the required assumptions, the test method is the independent (two-sample) t-test. In this case, we have:

- a large sample $(\mathbf{n_1} + \mathbf{n_2} > 100)$
- the population variance(s) are unknown, but we can assume = variances in 2 groups
- standard error of the means' difference is obtained as pooled estimate standard deviation of the sampling distribution of the difference

```
So we can compute: t_{calc} = \frac{\text{Difference Between Sample means}}{\text{Std. Err. of the difference}} = \frac{\bar{x_1} - \bar{x_2}}{\sqrt{\frac{s_1^2}{n_1} + \frac{s_2^2}{n_2}}}
```

```
1 # Step 1 - compute difference of sample means
2 mean_diff <- (mean_died - mean_survived) # -10.27645
3
4 # Step 2 - Compute associated t-statistics
5 # pooled std error
6 pooled_stderror <- sqrt(sd_died^2/(n_died ) + sd_survived^2/(n_survived ))
7 # pooled std error corrected
8 pooled_stderror_corr <- sqrt(sd_died^2/(n_died-1) + sd_survived^2/(n_survived-1))
9
10 ### t statistic
11 t_calc <- (mean_died - mean_survived) / pooled_stderror_corr</pre>
```

3.b Computation of the p-value associated to the t statistic

```
1 # Step 3 - degrees of freedom
2 # n1 + n2 - number of estimated parameters (2 means)
3 d_f <- n_died + n_survived - 1 - 1 # 297
4
5 # Step 4 - Deduced p-value
6 p_value <- 2 * pt(t_calc, df = d_f) # 0.4009635
7 p_value</pre>
```

[1] 0.4009635

4. Results and interpretation

1. Looking at the confidence interval of the difference, the sample mean_diff is well inside the 95% CI of = population mean

2. As for the p-value, $p_value = 0.40$ is bigger than threshold probability α

```
1 # Check
2 p_value

[1] 0.4009635

1 p_value < 0.05 # FALSE

[1] FALSE
```

DECISION: So, we fail to reject the null hypothesis of equal populations means of TPC. So the test indicates that we do not have sufficient evidence to say that the mean counts of platelets in between these two populations is different.

- EXAMPLE D -

(3+ samples | ANOVA test)

Comparing sample means from 3 or more groups (ANOVA)

In this example, we adopt the ANOVA ("Analysis Of Variance") test, i.e. an extension of the previous test, but examined how means of a variable differ across 3 or more groups. We will use 'one- way' ANOVA, which serves when there is only one explanatory variable ("treatment") with 3 or more levels, and only one level of treatment is applied for a given subject.

For this particular case, we use another realistic dataset showing the survival times of 33 laboratory mice with thymic leukemia who were randomly divided into 3 groups:

- 1st group received Treatment 1
- 2nd group received Treatment 2
- 3rd group as Control

1. Question: Is there a statistically significant difference between the mean values of the k po

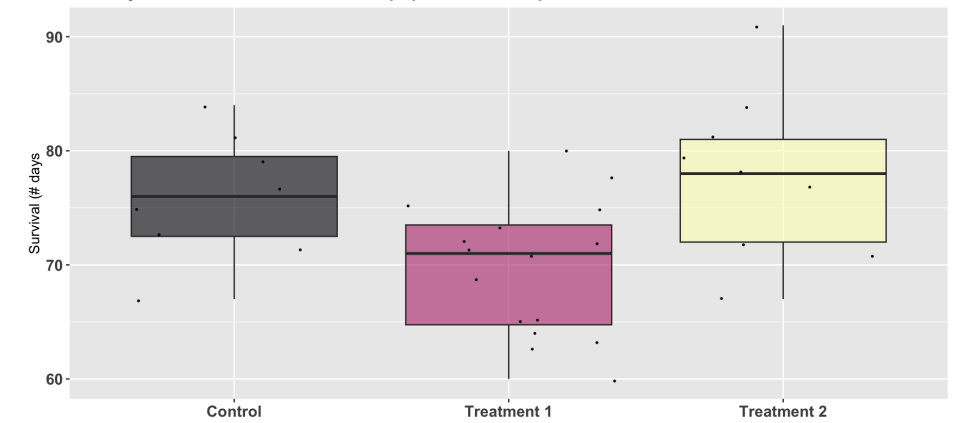
Defining the question formally:

- H_0 : $\mu_1 = \mu_2 = \mu_3$ all 3 population means are equal
- H_a : at least one of (μ_1, μ_2, μ_3) is not equal to the other means

1. Question: Is there a statistically significant difference between the mean values of the k po

The boxplot suggests that the 3 groups might have some fairly different distributions

Visually check mean and variance in populations' samples



2. Verify the assumptions for one-way ANOVA

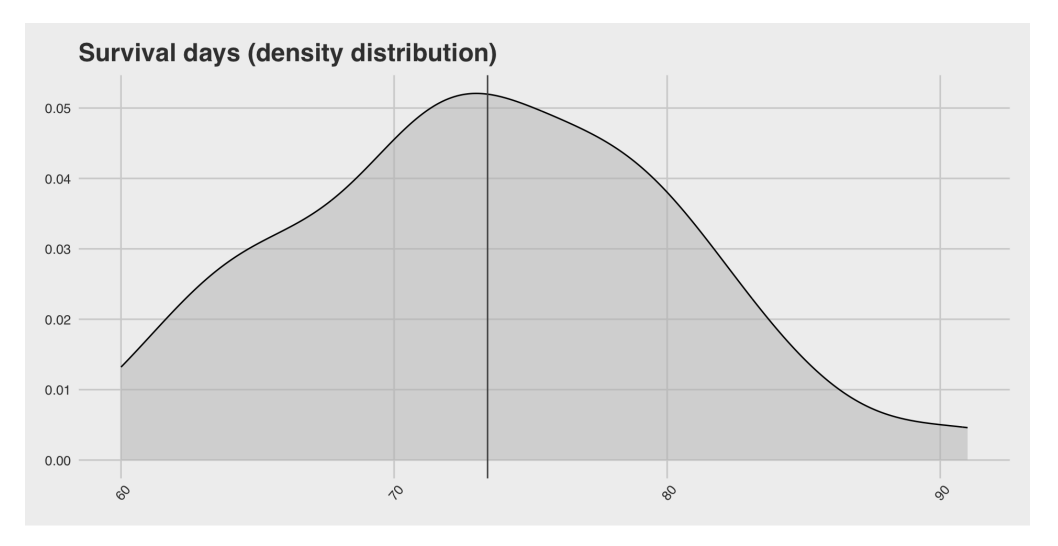
The dependent variable is on a metric scale. In the case of the analysis of variance, the independent variable (factor) has at least three levels.

Assumptions for the results of a one-way ANOVA to be valid:

- Independence of observations The observations in each group are independent of each other and the observations within groups were obtained by a random sample.
- 2. **Normally-distributed response variable** The values of the dependent variable follow a normal distribution. **?**
- 3. **Homogeneity of variance** The variances of the populations that the samples come from are equal. ?

Preliminary check for normality (visual)

2. Normally-distributed response variable (confirmed by visual inspection)



Preliminary check for normality (test) with stats::shapiro.test

```
3 stats::shapiro.test(mice[mice$group == "Control", "surv days", drop=TRUE])
   Shapiro-Wilk normality test
data: mice[mice$group == "Control", "surv days", drop = TRUE]
W = 0.99374, p-value = 0.9989
          1 stats::shapiro.test(mice[mice$group == "Treatment 1", "surv days", drop=TRUE])
   Shapiro-Wilk normality test
data: mice[mice$group == "Treatment 1", "surv days", drop = TRUE]
W = 0.95716, p-value = 0.6106
          1 stats::shapiro.test(mice[mice$group == "Treatment 2", "surv days", drop=TRUE])
    Shapiro-Wilk normality test
data: mice[mice$group == "Treatment 2", "surv days", drop = TRUE]
W = 0.97921, p-value = 0.9601
```

Preliminary check for normality (test) with rstatix::shapiro_test

(same thing, but using a different R function)

- 2. Normally-distributed response variable ✓
- (confirmed by Shapiro-Wilk normality test)

[The null hypothesis of this test is H_0 = "sample distribution is normal"]

```
1 # Shapiro-Wilk Normality Test to verify normality
2 # option 2 (all 3 groups at once)
3 mice %>%
4 dplyr::group_by(group) %>%
5 rstatix::shapiro_test(surv_days)
```

Preliminary check variance equality

3. Homogeneity of variance – ✓

(Besides visual inspection, confirmed by Levene test for variance equality)

[The null hypothesis H_0 = several groups have the same variance (possible variance differences occur only by chance, since there are small differences in each sampling)]

No evidence of violations of HOV were found, since the p-value for the Levene test (= 0.8427157) is greater than .05, then the variances are not significantly different from each other (i.e., the homogeneity assumption of the variance is met).

3 Computation of ANOVA F-ra

ANOVA in R can be done in several ways.

Since it's quite straightforward, let's do all the steps by hand first. We need to obtain the needed "ingredients" to calculate the F-ratio:

$$F_{\text{calc}} = \frac{\text{Mean Square Between}}{\text{Mean, Square Within}} = \frac{\text{MSB}}{\text{MSW}} = \frac{\frac{\text{SSB}}{\text{df 1}}}{\frac{\text{SSW}}{\text{df 2}}}$$

3.a Computation of ANOVA F-ratio ("by har

Option 1: Let's compute the ANOVA test "by hand"

```
2 mice calc <- mice %>%
     dplyr::mutate(mean all = mean(surv days),
            sd all = sd (surv days),
            dfw = 33-3, # df1 = n-k
            dfb = 3-1, # df2 = K-1
            group f = as.factor(group)
            dplyr::group by(group) %>%
     dplyr::mutate(n group = n(),
            mean group = mean(surv days),
            sd group = sd (surv days)) %>%
     ungroup() %>%
     mutate (ST = (surv days - mean all)^2,
             SW = (surv days - mean group)^2,
             SB = (mean group - mean all)^2
19 SST <- sum(mice calc$ST)</pre>
20 SSB <- sum(mice calc$SB)</pre>
21 SSW <- sum(mice calc$SW)
22 dfw <- 33-3 # df2
23 dfb <- 3-1 # df1
26 F calc <- (SSB/dfb)/(SSW/dfw) # 5.65
28 F crit <- qf(p = 0.01, df1 = 2, df2 = 30, lower.tail = FALSE) # 5.390346
```

3.b Computation of ANOVA F-ratio (with R function

That was just to show how to build it step-by-step (,), but we don't have to! We have alternative R functions that can do ANOVA for us:

Option 2: With the stats::aov followed by the command summary A

• Option 3: With the stats::oneway.test() function 🖺

```
One-way analysis of means

data: surv_days and group_f
F = 5.6522, num df = 2, denom df = 30, p-value = 0.008258
```

4. Results and interpretation

All 3 options have given the same results, i.e., F-ratio = 5.652 and a p-value = 0.00826

DECISION: Given that the p-value is smaller than 0.05, we reject the null hypothesis, so we reject the hypothesis that all means are equal. Therefore, we can conclude that *at least one* group is different than the others in mean number of survival days.



Note

Have you seen the kind of notation Pr(>F) 0.00826 ** before (as in the output of the stats::aov function)?

A CLOSER LOOK AT TESTING ASSUMPTIONS

- EXAMPLE E -

Testing two groups that are not independ

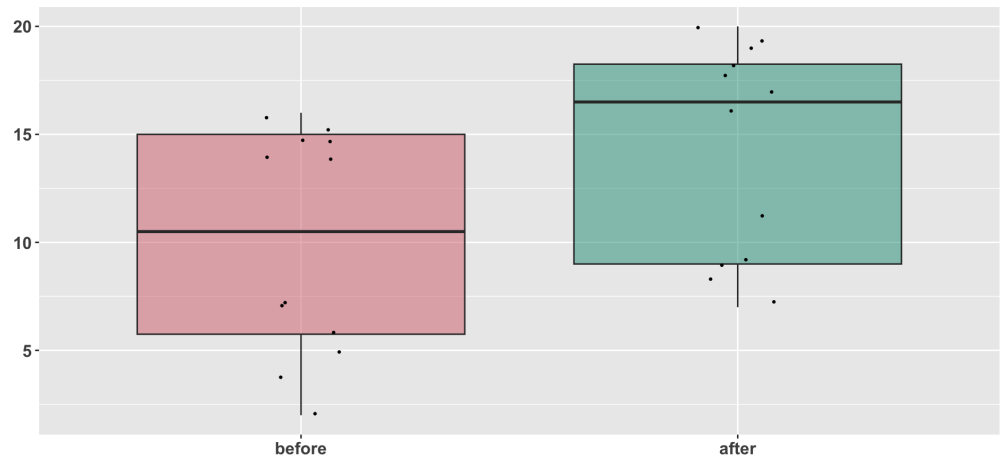
Let's introduce another toy dataset just for demonstration purposes: imagine a statistics test is administered to the *same* group of 12 students **before and after** attending a workshop .

```
1 # toy dataset for paired groups
2 grades <- data.frame(
3 before = c(16, 5, 15, 2, 14, 15, 4, 7, 15, 6, 7, 14),
4 after = c(19, 18, 9, 17, 8, 7, 16, 19, 20, 9, 11, 18)
5 )</pre>
```

We may reshape the dataframe into the long form using tidyr::pivot_longer (for plotting)

1. Question: Is the difference between two PAIRED samples statistically signif

Boxplot of test grades grouped as before and after



What a successful workshop!

2 Hypotehsis for the PAIRED t-test for dependent sam

In this example, it is clear that the two samples are not independent since the same 12 students took the test before and after the workshop.

Given that the normality assumption is NOT violated (and given the small sample size), we use the **paired t-test**, with the following hypotheses:

- ullet H_0 : mean grades before and after the workshop are equal
- ullet H_a : mean grades before and after the workshop are different

2 Computation of the PAIRED t-test for dependent sam

```
1 t stat paired <- stats::t.test(x = grades$before,</pre>
                                             y = grades$after,
                                             mu = 0,
                                             alternative = "two.sided",
                                             paired = TRUE
          7 t stat paired
    Paired t-test
data: grades$before and grades$after
t = -1.8777, df = 11, p-value = 0.08718
alternative hypothesis: true mean difference is not equal to 0
95 percent confidence interval:
-9.2317713 0.7317713
sample estimates:
mean difference
          -4.25
          2 t calc pair <- t stat paired[["statistic"]][["t"]] # -1.877683</pre>
```

3 p value pair <- t stat paired[["p.value"]] # 0.08717703</pre>

3. Results and interpretation

We obtain the test statistic, the p-value and a reminder of the hypothesis tested.

The calculated **t value** is -1.8776829 The **p-value** is 0.087177. Therefore, at the 5% significance level, **we do not reject the null hypothesis** that the statistics' grades are similar before and after the workshop (**6**).

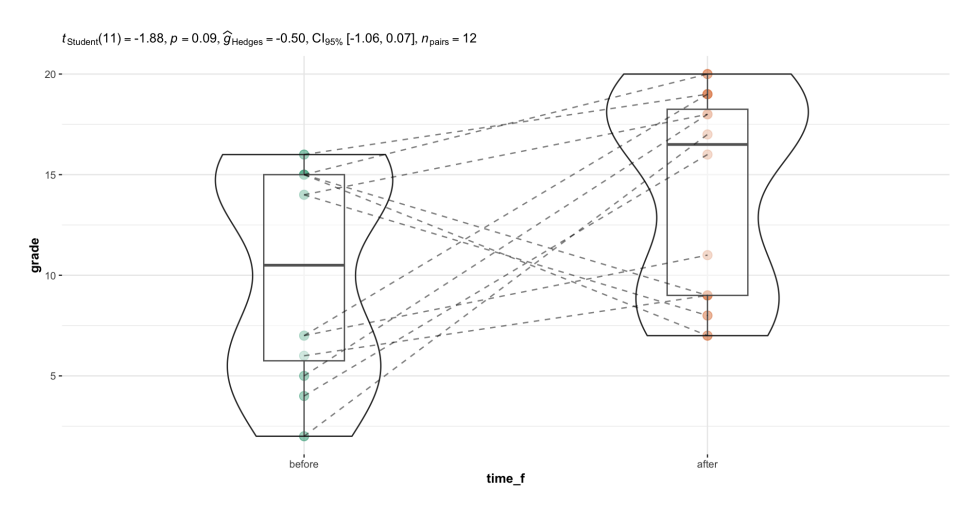
Bonus function!

It is worth mentioning the ggstatsplot package, which combines plots representing the distribution for each group—and the results of the statistical test displayed in the subtitle of the plot.

Below we check out the ggwithinstats() function for paired samples.

Bonus function!

The test results are rendered with the plot!



— EXAMPLE F —

(2 samples no normal | Wilcoxon Rank Sum Test)

Testing samples without normality assumption

Let's go back to the HEART FAILURE dataset but looking at the levels of **Creatinine Phosphokinase (CPK)** in the blood, an enzyme that might indicate a heart failure or injury

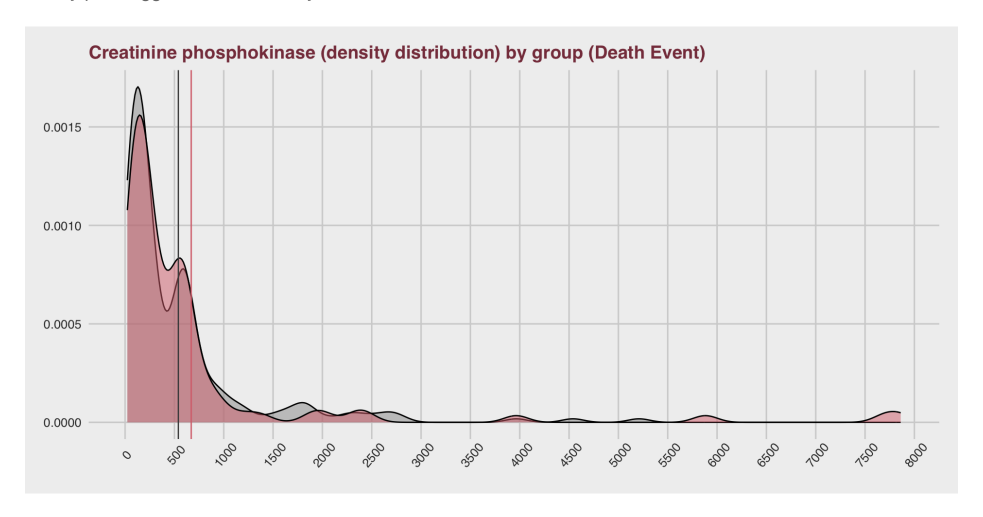
1. Question: Is there a statistically significant difference between CPK levels in the blood of

Defining the question formally:

- H_0 : $\mu_{\rm CPK-died}=\mu_{\rm CPK-surv}$ there is no difference in mean CPK between patients who suffered heart failure and died versus patients who survived after heart failure
- H_a : $\mu_{\text{CPK-died}} \neq \mu_{\text{CPK-surv}}$ there is a difference in mean CPK between patients who suffered heart failure and died versus patients who survived after heart failure (two-sided test)

1. Question: Is there a statistically significant difference between CPK levels in the blood of

The density plot suggests non normality of the variable distribution



Preliminary check for normality (visual)

Normally-distributed response variable - X

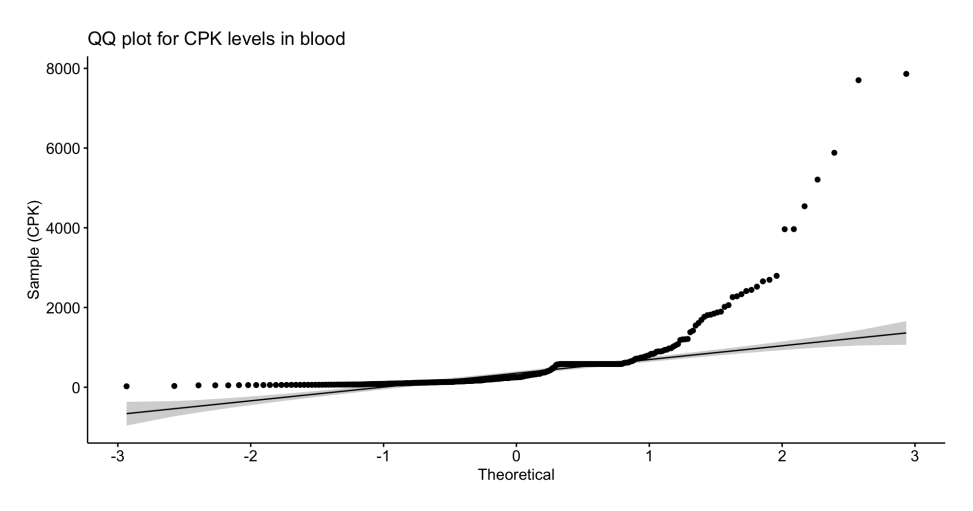
QQ plot (or quantile-quantile plot) draws the correlation between a given sample and the normal distribution. A 45-degree reference line is also plotted. In a QQ plot, each observation is plotted as a single dot.

• If the data are normal, the dots should form a straight line.

```
1 # visual verification with QQ plot
2 ggpubr::ggqqplot(
3 heart_failure$creatinine_phosphokinase,
4 title = "QQ plot for CPK levels in blood",
5 xlab ="Theoretical", ylab = "Sample (CPK)")
```

Preliminary check for normality (visual)

In a QQ plot, if the data are normal, the dots should follow a straight line.



Preliminary check for normality (test) with rstatix::shapiro_test

(same thing, but using a different R function)

- Normally-distributed response variable X
 - (NOT normality confirmed by Shapiro-Wilk normality test)

[The null hypothesis of this test is H_0 = "sample distribution(s) is/are normal"] Given the p-value we reject the null hypothesis

```
2 heart failure %>%
              dplyr::group by(DEATH EVENT f) %>%
              rstatix::shapiro test(creatinine phosphokinase)
# A tibble: 2 \times 4
 DEATH EVENT f variable
                                        statistic
                                            <dbl>
  <fct>
               <chr>
                                                     <dbl>
1 survived
               creatinine phosphokinase
                                            0.628 8.51e-21
               creatinine phosphokinase
2 died
                                            0.439 1.99e-17
```

3. Computation of the Wilcoxon Rank Sum test stat

The Wilcoxon Rank Sum test is considered to be the nonparametric equivalent to the two-sample independent t-test

Its ASSUMPTIONS are:

- Ordinal or Continuous dependent variable: e.g. CPK levels
- Independence: All of the observations from both groups are independent of each other **V**
- Shape: The shapes of the distributions for the two groups are roughly the same

```
wrs res <- wilcox.test(creatinine phosphokinase ~ DEATH EVENT, # immagino 0, 1
                     data = heart failure ,
                     exact = FALSE,
                     alternative = "two.sided" )
5 wrs res
```

Wilcoxon rank sum test with continuity correction

```
data: creatinine phosphokinase by DEATH EVENT
W = 9460, p-value = 0.684
alternative hypothesis: true location shift is not equal to 0
```

4. Results and interpretation

RESULTS: since the test statistic is W = 9460 and the corresponding p-value is 0.684 > 0.05, we fail to reject the null hypothesis.

INTERPRETATION: We do not have sufficient evidence to say that CPK levels for dead patients is different than that of survived patients $\mu_{\text{CPK-died}} \neq \mu_{\text{CPK-surv}}$ at some statistically significant level)

- EXAMPLE G -

(2 samples no HOV I t test with the Welch correction)

Testing samples without homogeneous variance of observations assumption

1. Question: Is there a statistically significant difference between serum sodium levels in the

Defining the question formally:

- H_0 : $\mu_{
 m sersod-died}=\mu_{
 m sersod-surv}$ there is no difference in mean serum sodium between patients who suffered heart failure and died versus patients who survived after heart failure
- H_a : $\mu_{\text{sersod-died}} \neq \mu_{\text{sersod-surv}}$ there is a difference in mean serum sodium between patients who suffered heart failure and died versus patients who survived after heart failure (two-sided test)

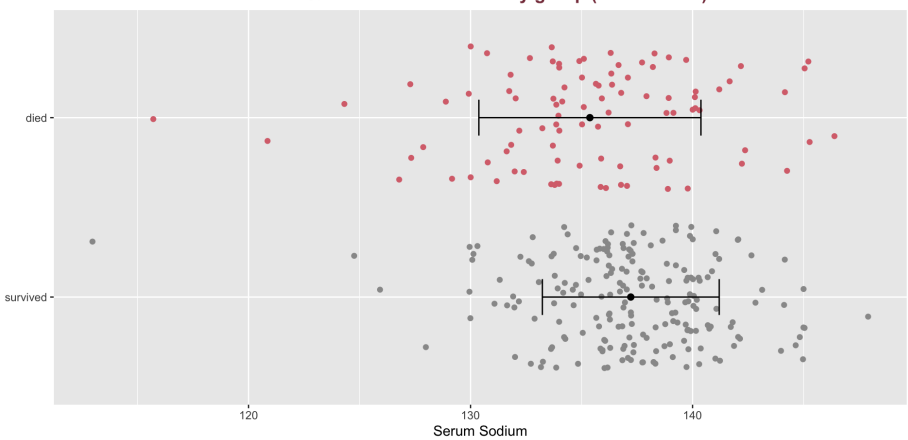
Preliminary check "HOV" assumption (visual)

• Homogeneity of Variance assumption - X Plotting the data offers some graphical intuition that the variance of observations in the two groups seem not homogenous

```
2 swstats <- heart failure %>%
     group by (DEATH EVENT f) %>%
     summarise(count = n(),
     mean = mean(serum sodium,na.rm=TRUE),
      stddev = sd(serum sodium, na.rm=TRUE),
      meansd l = mean - stddev,
       meansd u = mean + stddev)
11 ggplot(swstats, aes(x=DEATH EVENT f, y=mean)) +
     geom point(colour = "black" , size = 2) +
     geom point(data=heart failure, aes(x=DEATH EVENT f, y=serum sodium, colour = DEATH EVENT f),
                position = position jitter() ) +
     scale colour manual(values = c("#9999999","#d8717b") ) +
     geom errorbar(aes(ymin = meansd 1, ymax = meansd u), width=0.2, color = "black") +
     labs(title = "Mean (-/+SD) serum sodium (mEq/L) by group", x = "", y = "Serum Sodium") +
     guides(fill = "none") +
     coord flip() +
     labs(title = "Serum Sodium means and 95% confidence intervals by group (Death Event)") +
     theme(legend.position="none",plot.title = element text(size = 14,face="bold", color = "#873c4a"))
```

Preliminary check "HOV" assumption (visual)

Serum Sodium means and 95% confidence intervals by group (Death Event)



Preliminary check "HOV" assumption (test)

It is always best to use an actual test, so we use also the **Fisher's F test** to verify equal variances of Serum Sodium concentration in the two groups. [In this test H_0 = "the ratio of variances is equal to 1"]

F test to compare two variances

```
data: heart_failure$serum_sodium[heart_failure$DEATH_EVENT == 1] and
heart_failure$serum_sodium[heart_failure$DEATH_EVENT == 0]
F = 1.5769, num df = 95, denom df = 202, p-value = 0.007646
alternative hypothesis: true ratio of variances is not equal to 1
95 percent confidence interval:
    1.127401 2.254466
sample estimates:
ratio of variances
    1.576922
```

Given the p-value = 0.007646 (smaller than α) we reject the null hypothesis, hence the HOV assumption for the t test does not hold.

2 Computation of the t test with the Welch correc

We can still run the **t test but with Welch correction**, i.e. the unequal variance condition is compensated by lowering the df. In fact the documentation (?t.test), reads:

- If var.equal = TRUE, then the pooled variance is used to estimate the variance
- Otherwise (var equal = FALSE), the Welch approximation to the degrees of freedom is used.

Welch Two Sample t-test

```
data: heart_failure$serum_sodium[heart_failure$DEATH_EVENT == 1] and
heart_failure$serum_sodium[heart_failure$DEATH_EVENT == 0]
t = -3.1645, df = 154.01, p-value = 0.001872
alternative hypothesis: true difference in means is not equal to 0
95 percent confidence interval:
    -2.9914879 -0.6920096
sample estimates:
mean of x mean of y
135.3750 137.2167
```

3. Results and interpretation

RESULTS: since the test statistic is t = -3.1645 (with df = 154.01) and the corresponding p-value is 0.001872 < 0.05, we reject the null hypothesis.

INTERPRETATION: We therefore have sufficient evidence to say that the level of serum sodium levels for dead patients is significantly different than that of survived patients $\mu_{\text{sersod-died}} \neq \mu_{\text{sersod-surv}}$

Final thoughts/recommendations

- There are often many ways to do the same thing in R (which is both a blessing and a curse in open source software). Which should you choose? It depends on the situation, but you may want to consider:
 - how recent/popular/well maintained is a {package} (this affects its stability)
 - the more a function abstracts away complexity, the easier it is to use interactively, but the harder it gets to handle inside your own custom functions
 - different function outputs may be more/less suitable for your analysis/publication requirements (check out your peers' choices!)
 - (Always read the documentation to assess all of the above)
- With easy equations, breaking them down "by hand" (at least once!) can really help you understand them
- It may seem a lot of work to write R code the first time (e.g. for a publication-ready plot), but the good news is **once you wrote a script, you will be able to easily re-use it in many more instances** (**) (**)
- Sample size n has a very powerful impact on classical hypothesis testing results! More on this later...