



# **Data Analysis with R:**

Lecture Slides (all)

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October 2018

#### Goals of the course

#### To be able to...

- import data sets to R
- describe data with R
- apply basic statistical tests in R
- some ideas for more advanced statistical tools ...
- simulate a data set similar to own research

#### **General remarks**

#### Course schedule:

- Starting at 9:00am / 9:30am (?)
- Tea breaks in between
- Lunch break
- Teaching until 4.30pm ( $\sim$  5pm)

#### Optaining a certificate is conditional on:

- active participation in class
- attending at least 75 % of the course (lecture & exercises)
- short final exam (format to be defined)

#### Getting to know each other

- My name is ...
- I am doing a Master / a PhD in ...
- I hope to learn in this course how to ....
- My personal goal for this course is ...

# How do we reach these goals

- hands on exercises with R:
  - chickwts
  - ToothGrowth
  - bacteria
  - perulung
  - ... and others.
- interactive discussions & student's present their own solutions
- ask us a lot of questions but also ask google for help!
- group work
- short motivational lectures

#### Get started with data set: chickwts

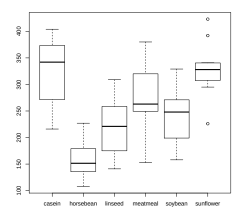
An experiment was conducted to measure and compare the effectiveness of various feed supplements on the growth rate of chickens.

```
# load data set "chickuts"
data(chickwts)
# the head(...) function shows the first 6 observations
head(chickwts)
    weight
           feed
## 1 179 horsebean
## 2 160 horsebean
## 3 136 horsebean
## 4 227 horsebean
## 5 217 horsebean
## 6 168 horsebean
# FUNCTION - open bracket - DATA SET / VARIABLE - close bracket
```



# Ideas for plotting the data

```
# use x axis to show the categorical variable (feed),
# y axis to represent the continuous variable (weight)
# boxplot (y.cont.variable ~ x.cat.variable, data = dataset)
# ?boxplot
boxplot(weight ~ feed, data = chickwts)
```





#### Ideas for analysing the data

```
anova <- aov(weight ~ feed, data = chickwts)
summarv(anova)
## Df Sum Sq Mean Sq F value Pr(>F)
## feed 5 231129 46226 15.37 5.94e-10 ***
## Residuals 65 195556 3009
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(aov(weight ~ feed, data = chickwts))
           Df Sum Sq Mean Sq F value Pr(>F)
##
## feed 5 231129 46226 15.37 5.94e-10 ***
## Residuals 65 195556 3009
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

# Functionalities in R and RStudio A hands on example

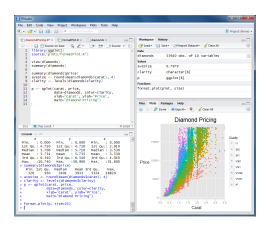


```
x <- c(0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10)
y <- c(20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30)
plot(x, y)
```

#### **Functionalities in R and RStudio**



- Source
- Console
- Environment, History, Files
- Files, Plots, Packages, Help



# Good housekeeping!



- Define manually a new folder called RCourse in your personal documents on your personal computer
- Know in which directory you are

```
getwd()
## [1] "/home/mburi/ownCloud/git/DataAnalysisWithR/Lectures"
```

Set directory path

```
# back- and forslash is dependent on the system
setwd("C:/Users/muriel/Documents/RCourse/")
setwd("C:\\Users\\muriel\\Documents\\RCourse\\")
```

Always clean up before starting with new R-Script

```
rm(list=ls()) # empty workspace, delete previously saved variables
```

# How to get help in R



?chickwts
?boxplot

Also, have a look at the examples at the end of the help pages.

#### **Exercise 1**



#### A data frame in R: chickwts



# chickwts[ ROWS , COLUMNS ]



chickwts[6,1]

	weight <sup>°</sup>	teed
1	179	horsebean
2	160	horsebean
3	136	horsebean
4	227	horsebean
5	217	horsebean
6	168	horsebean
7	108	horsebean
8	124	horsebean
9	143	horsebean
10	140	horsebean
11	309	linseed
12	229	linseed
13	181	linseed

chickwts[11, 2]

#### Rows and columns of a data frame: chickwts



#### Values of ...

```
# ... all columns of sixth observation:
chickwts[6,]
# ... all columns of sixth to eleventh observation:
chickwts[c(6:11), ]
# ... all columns of sixth, eleventh and twentieth observation:
chickwts[c(6, 11, 20), ]
# ... all rows of first column (weight):
chickwts[ . 1]
# ... all rows of second column (feed):
chickwts[ . 2]
# or use the "$" sign as a reference to column "feed":
chickwts$feed
```

#### What is a data frame in R?



A data frame is used for storing a list of vectors of equal length. For example, the following variable df is a data frame containing three vectors n, s, b.

```
n <- c(2, 3, 5)
s <- c("aa", "bb", "cc")
b <- c(TRUE, FALSE, TRUE)
df <- data.frame(n, s, b) # df is a data frame</pre>
```

#### The characteristics of a data frame are:

- The column names should be non-empty.
- The row names should be unique.
- Each column should contain same number of data items.

#### Data frame in R



```
a \leftarrow c(1, 2, 3, 4)
а
## [1] 1 2 3 4
data.frame(a)
## a
## 1 1
## 2 2
## 3 3
## 4 4
b <- c("d", "h", "h", "d")
dat <- data.frame(a, b)</pre>
dat
## a b
## 1 1 d
## 2 2 h
## 3 3 h
## 4 4 d
```

#### Data frame in R: How to add a variable (var)

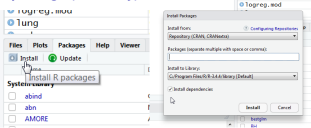


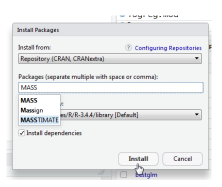
```
my.var \leftarrow c(1.3, 1.5, 1.8, 2.4)
# use "$" to refer to the additional vector variable
dat$my.var1 <- my.var</pre>
dat$my.var2 <- my.var</pre>
dat
## a b my.var1 my.var2
## 1 1 d 1.3 1.3
## 2 2 h 1.5 1.5
## 3 3 h 1.8 1.8
## 4 4 d 2.4 2.4
# What is the dimension (number of rows and columns) of our data frame?
dim(dat) # 4 rows and 4 columns
## [1] 4 4
```

#### **Exercise 2**



# How to install a package (manually) in R





# Using R is like cooking ...

Get into the kitchen	Change working directory
Get specialist electric tools into your kitchen (e.g. blender, ice- cream maker, etc.)	Install packages
Switch on your specialist electric tools	Load packages using the "library" function
Bring in your ingredients	Import data and save to R data frames
Check your ingredients	Use the function "summary" and basic tables to check your data for missing or implausible values (e.g. a number in a variable where "yes" or "no" are expected
Chop things up (if required)	Split or filter data
Cook, using general and specialist tools	Carry out further descriptive and test statistics

# How to install a package in R



```
# INSTALL package (only done ONCE!)
install.packages("MASS")
# LOAD package (whenever you use something from it!)
library("MASS")
data(bacteria)
?bacteria
```

#### **Exercise 3**





• Google for select observations in R.

# Creating and assigning objects in R



Objects are assigned values using <-, an arrow formed out of < and -. For example, the following command assigns the value 1 to the object a.

```
a <- 1 # ALWAYS use "gets" assignment operator!
# a = 1 # DO NOT USE the equal sign as the assignment operator!
```

After this assignment, the object a contains the value 1. Another assignment to the same object will change its value.

a <- 5

# **Examples of assigned objects: single number**



```
a <- 1
b <- 2
c <- a + b # c = 3
c
```

# **Examples of assigned objects: vector**



```
a <- c(1, 2, 3, 4, 5)
b <- 1
c <- a + b
c
## [1] 2 3 4 5 6
```

# Examples of assigned objects: model



```
anova_model <- aov(weight ~ feed, data = chickwts)
summary(anova_model)

## Df Sum Sq Mean Sq F value Pr(>F)

## feed 5 231129 46226 15.37 5.94e-10 ***

## Residuals 65 195556 3009

## ---

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

#### Examples of assigned objects: data frame



```
bac <- bacteria
str(bac) # $ week: int 0 2 4 11 0 2 6 11 0 2 ...
## 'data frame': 220 obs. of 6 variables:
##
   $ y : Factor w/ 2 levels "n", "y": 2 2 2 2 2 2 1 2 2 2 ...
   $ ap : Factor w/ 2 levels "a", "p": 2 2 2 2 1 1 1 1 1 1 1 ...
##
##
   $ hilo: Factor w/ 2 levels "hi","lo": 1 1 1 1 1 1 1 1 2 2 ...
##
   $ week: int 0 2 4 11 0 2 6 11 0 2 ...
## $ ID : Factor w/ 50 levels "X01","X02","X03",..: 1 1 1 1 2 2 2 2 3 3 ...
##
   $ trt : Factor w/ 3 levels "placebo", "drug",...: 1 1 1 1 3 3 3 3 2 2 ...
bac sub <- subset(bac, week == 2)
str(bac_sub) # $ week: int 2 2 2 2 2 2 2 2 2 2 ...
## 'data.frame': 44 obs. of 6 variables:
   $ y : Factor w/ 2 levels "n", "y": 2 2 2 2 2 2 1 2 2 2 ...
##
   $ ap : Factor w/ 2 levels "a","p": 2 1 1 2 2 1 1 2 2 2 ...
##
   $ hilo: Factor w/ 2 levels "hi","lo": 1 1 2 2 2 2 1 1 2 1 ...
##
   $ week: int 2 2 2 2 2 2 2 2 2 2 ...
##
## $ ID : Factor w/ 50 levels "X01", "X02", "X03", ...: 1 2 3 4 5 6 7 8 9 11 ...
   $ trt : Factor w/ 3 levels "placebo", "drug",..: 1 3 2 1 1 2 3 1 1 1 ...
```

#### Structure of a R objects



The str function displays the structure of an R object. One line for each "basic" structure is displayed.

```
## 'data.frame': 44 obs. of 6 variables:
## $ y : Factor w/ 2 levels "n","y": 2 2 2 2 2 2 1 2 2 2 ...
## $ ap : Factor w/ 2 levels "a","p": 2 1 1 2 2 1 1 2 2 2 ...
## $ hilo: Factor w/ 2 levels "hi","lo": 1 1 2 2 2 2 1 1 2 1 ...
## $ week: int 2 2 2 2 2 2 2 2 2 2 2 ...
## $ ID : Factor w/ 50 levels "X01","X02","X03",..: 1 2 3 4 5 6 7 8 9 11 ...
## $ trt : Factor w/ 3 levels "placebo","drug",..: 1 3 2 1 1 2 3 1 1 1 ...
```

#### **Exercise 4**



#### Data types in R

#### numeric



```
data(ToothGrowth)
ToothGrowth$len[1:6]
## [1] 4.2 11.5 7.3 5.8 6.4 10.0
class(ToothGrowth$len[1:6])
## [1] "numeric"
```

#### integers

```
bacteria$week[1:6]
## [1] 0 2 4 11 0 2

class(bacteria$week[1:6])
## [1] "integer"
```

# (un/ordered) factor

```
chickwts$feed[1:6]
## [1] horsebean horsebean horsebean horsebean horsebean
## Levels: casein horsebean linseed meatmeal soybean sunflower
levels(chickwts$feed)[1:3]
## [1] "casein" "horsebean" "linseed"
```

# **Data types in R: Ordered Factors**



#### Ordinal variables are represented as ordered factors:

```
bac_growth <- c("none", "+", "++", "+", "+++", "+", "none") # vector
bac growth <- factor(bac growth, levels = c("none", "+", "++", "+++"),
                    order = TRUE)
bac_growth
## [1] none + ++ + +++ +
                                   none
## Levels: none < + < ++ < +++
mood <- c("OK", "Well", "Super", "Super", "Don't ask", "OK") # vector</pre>
mood <- factor(mood, levels = c("Don't ask", "Well", "OK", "Super"),</pre>
              order = TRUE)
mood
## [1] OK Well Super Super Don't ask OK
## Levels: Don't ask < Well < OK < Super
```

#### **Exercise 5**



#### **Exercise 6**



# **Examples of different data types**

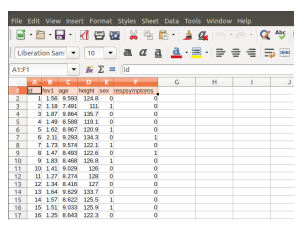


- numeric variable
- integer variable
- variable with two levels (binary factor)
- ordered variable with more than two levels (ordinal)
- unordered variable with more than two levels (nominal)

## Rules for importing data into R (from Excel)



- First row of excel sheet contains variable names:
   y, ap, hilo, week, ID, trt.
- Columns of excel sheet represent variables.
- Rows of excel sheet represent observations per individual (except for the first row).



### Rules for naming variables

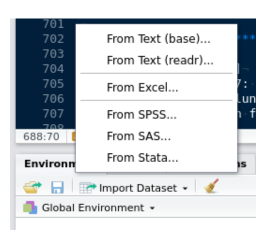


#### Variable names should ...

- start with a letter (not a number):
   y, ap, hilo, week, ID, trt
- longer variables names should be separated with dots:
   time.in.weeks
- do not use special characters, such as /, #, @, &, ★, ...

#### How to import external data files into R?

- > Import Dataset > From Text (base)... > CSV Files (.csv) or
- > Import Dataset > From Excel...



## How to import external data files into R?



- Environment (upper right corner)
- > Import Dataset > From Text (base)... > CSV Files (.csv)

- Import Dataset > From Text (base)... > Text Files (.txt)
- > Import Dataset > From Excel... > Excel Files (.xlsx)

```
install.packages("readxl")
library("readxl")
perulung_ems <- read_excel("perulung_ems.xlsx")
lung <- data.frame(perulung_ems)
head(lung)</pre>
```

## How to import .txt and .csv files into R? (1/2)

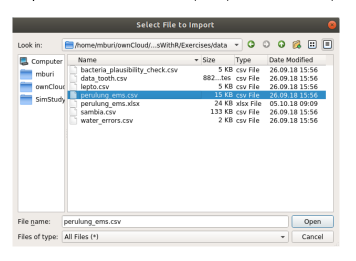


- Environment (upper right corner)
- > Import Dataset > From Text (base)... > CSV Files (.csv)

### How to import .txt and .csv files into R? (1/2)

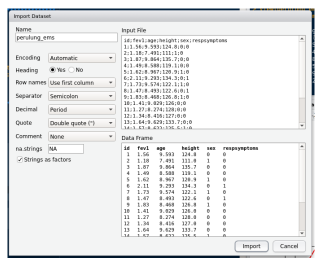


- Environment (upper right corner)
- > Import Dataset > From Text (base)... > CSV Files (.csv)



#### How to import .txt and .csv files into R? (2/2)





# How to import .xlsx files into R? (1/3)

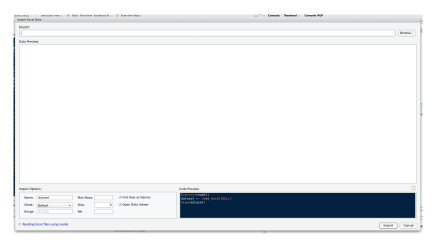


- Environment (upper right corner)
- > Import Dataset > From Excel... > Excel Files (.xlsx)

# How to import .xlsx files into R? (1/3)

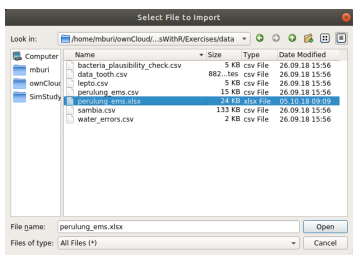


- Environment (upper right corner)
- > Import Dataset > From Excel... > Excel Files (.xlsx)



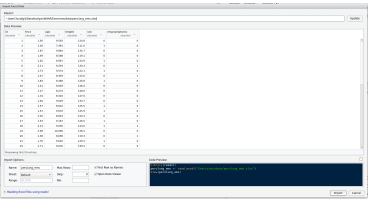
#### How to import .xlsx files into R? (2/3)





#### How to import .xlsx files into R? (3/3)





```
perulung_ems <- read_excel("perulung_ems.xlsx")
lung <- data.frame(perulung_ems)
head(lung)</pre>
```

#### **Exercise 7: perulung**



Data from a study of lung function among children living in a deprived suburb of Lima, Peru. Data taken from Kirkwood and Sterne, 2nd edition.

#### Variables:

- fev1: in liter, "forced expiratory volume in 1 second" measured by a spirometer. This is the maximum volume of air which the children could breath out in 1 second
- age: in years
- height: in cm
- sex: 0 = girl, 1 = boy
- respsymp: respiratory symptoms experienced by the child over the previous 12 months

#### Why do we need Statistics?

# Repeatability of results:

**Statistical science** allows us to estimate what might happen if an experiment was repeated - but without having to actually repeat it!

#### Why do we need Statistics?

- Study results must be shown to be robust, i.e. real and not due to random chance
- Best way to demonstrate this is to repeat the same experiment/study many times each with different subjects (animals) drawn from the same study population and show that the result is truly repeatable
- It is generally totally impractical, in terms of both time and resources, to repeat an experiment many times!

#### Why do we need Statistics?

- Instead of repeating the experiment many times probability theory i.e. statistics is used to estimate what might have happened if the experiment had been repeated
- A mathematical model is used to fill this "data gap"
- Generally the most difficult task in statistics is to decide what "model" is most appropriate for a given experiment

#### What is Statistics? - A definition

A set of analytical tools designed to quantify uncertainty

- If an experiment or procedure is repeated, how likely is it that the new results will be similar to those already observed?
- What is the likely variation in results if the experiment was repeated?

#### What is Statistics? - A definition

#### The key scientific purpose of statistics

- to provide evidence of the existence of some "effect" of scientific interest
- i.e. evidence based medicine

#### As a reminder: The importance of study design

Even the most sophisticated statistical analyses cannot rescue a poorly designed study

- → unreliable results
- ightarrow inability to answer the main research question

#### **Putting Statistics in Context**

- The vast majority of analyses can be done in a straightforward fashion - just remember and always use common sense as a guide - be skeptical!
- It is very easy to get "lost" in the statistical software and technical jargon, which differs markedly between different software packages. Terminology can also differ greatly between textbooks...
- Wikipedia is as good a resource as any for finding out about different statistical tests and terminology

 It is crucially important to explore your data fully before considering any "formal" statistical analyses

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- What explorations are done depends on the objective of the study - the research question(s)
- Helps to decide what kind of formal statistical analyses might be most appropriate for the data available
- What a simple descriptive analysis does not provide is evidence of whether the observed treatment effects are large enough to be notable once sampling variation has been accounted - that is the role of formal analyses, e.g. hypothesis testing

# **Summary Statistics Continuous (Integers / Numeric)**

- Mean a measure of location. Always examine the average value of the response variable(s) for the different "treatment" effects in your data
- Median a robust single value summary of a set of data (50% quantile point) - most useful in highly skewed data or data with outliers
- Standard deviation (sd) a measure of spread, how variable the data are
- Standard error of the mean (se) an estimate of how far the sample mean is likely to be from the population mean
- and others: min, max, range, IQR, ...

# **Continuous (Integers / Numeric) Summary Statistics**



```
mean(x) # mean
median(x) # median
sd(x) # standard deviation
min(x) # minimum
max(x) # maximum
range(x) # range
IQR(x) # interquartile range
```

#### **Continuous Data Summaries**

#### standard deviation

$$s = \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2}$$

#### standard error

$$se = \frac{s}{\sqrt{n}}$$

# Correlation coefficient Combination of continuous and continuous

Correlation coefficient a measure association between two continuous variables (common but somewhat limited)

#### Pearson's correlation coefficient r

$$\mathsf{r} \! = \! \frac{\sum_{i=1}^{n} (X_i \! - \! \bar{X}) (Y_i \! - \! \bar{Y})}{\sqrt{\sum_{i=1}^{n} (X_i \! - \! \bar{X})^2} \sqrt{\sum_{i=1}^{n} (Y_i \! - \! \bar{Y})^2}}$$

 $\bar{X}$ : mean of variable x

 $\overline{Y}$ : mean of variable y

#### Correlation of continuous and ordinal variables



```
# Test for Association/Correlation Between
# Paired Samples
cor.test(data$x, data$y, method = "pearson")
cor.test(data$x, data$y, method = "spearman")
# Scatterplot(s)
pairs(data$x ~ data$y)
pairs(data)
```

# Summary Statistics Continuous and ordinal variables



```
tapply(data$x.cont, data$y.fac, mean)
tapply(data$x.cont, data$y.fac, median)
tapply(data$x.cont, data$y.fac, sd)
```

# Summary Statistics Ordinal



- Median a robust single value summary of a set of data (50% quantile point) most useful in highly skewed data or data with outliers
- e.g.10th and 90th percentile a measure of spread, how variable the data are

```
quantile(x, probs = c(0.1, 0.9))
```

proportions - e.g. percentage per grade

```
prop.table(table(data$x.fac))
prop.table(table(data$x.fac, data$y.fac))
```

# **Summary Statistics Nominal**



- proportions percentage within the different categories
- contingency tables e. g. 2 x 2

```
table(data$x.fac)
table(data$x.fac, data$y.fac)
prop.table(table(data$x.fac))
prop.table(table(data$x.fac, data$y.fac))
```

#### **Exercise 8**



#### How to deal with missing values in R? (1/3)

- In R, missing values are represented by the symbol NA (not available).
- Impossible values (e. g., dividing by zero) are represented by the symbol NaN (not a number).
- Ask yourself why a NA and / or NaN occurs!

#### How to deal with missing values in R? (2/3)

### Testing for Missing Values

```
vec1 <- c(1, 2, 3, NA)
is.na(vec1) # returns a vector (FALSE, FALSE, FALSE, TRUE)
# The TRUE indicates the position of the NA in vec1.</pre>
```

## Recoding Values to Missing

```
# recode specific values (e. g. 0.001) to missing for variable x # select rows where x is 0.001 and recode value in column x with NA tmp.row <- which(dat$x == 0.001) dat$x[tmp.row] <- NA
```

## How to deal with missing values in R? (3/3)

Excluding Missing Values from specific function calls

```
a <- c(1, 2, NA, 3)
mean(a) # returns NA
mean(a, na.rm=TRUE) # returns 2
```

 Check for complete cases with function complete.cases(...)

```
# list rows of data that have missing values
dat[!complete.cases(dat),]
subdat <- dat[complete.cases(dat),]</pre>
```

 Create new dataset without missing data with function na.omit(...)

```
new.dat <- na.omit(dat)
```

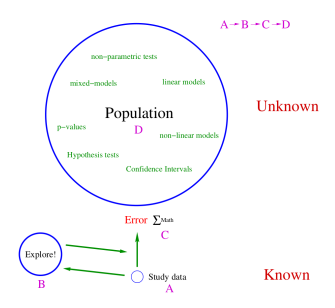
## How to check your data for plausibility?

- Ask yourself what can go wrong?
- Implausible values?
- Impossible values?
- Logical errors?

# Exercise 9A and 9B Plausibility Checks & Missing Values

## **Exercise 10**

#### **Overview**



#### **Basic Statistical Tests**

Study data is collected for a purpose - to answer one or more specific scientific questions. The classical way to perform a formal statistical analyses of these data is to formulate these research questions into statistical **hypothesis tests**.

In this section we will go through a simple example in detail to highlight some of the important concepts - the general approach for more complex analyses is exactly same. *Note: the precise technical details are much less important than the concepts!* 

## After six weeks will the mean weight of a chicken be more than 250 grams?

There are 71 observations in chickwts from which to answer this question. This can be formulated into a statistical hypothesis test. A hypothesis test has two parts, the null hypothesis and the alternative hypothesis. This is typically written as follows:

 $H_0: \mu \le 250$  $H_A: \mu > 250$ 

where  $\mu$  is the mean weight in the **population** of chickens from which the sample of 71 chickens was drawn. Remember - we know the mean weight in the sample of chickens is greater than 250 it is the **population** of chickens which we are interested in.

## After six weeks will the mean weight of a chicken be at least 250 grams?

 $H_0: \mu \le 250$  $H_A: \mu > 250$ 

The null hypothesis  $(H_0)$  is the default situation, sometimes called the "state of nature". In a treatment-control trial,  $H_0$  is typically that the effect of the treatment is not different from the control. In this example our default position is that the mean weight of chickens is  $\leq$  250. This is called a single-sided hypothesis test.



We now analyse the 71 observations to see whether there is evidence to **REJECT** the null hypothesis  $H_0$ , and if the null hypothesis is rejected then we can conclude that the available evidence supports the alternative hypothesis.

```
t.test(chickwts$weight, mu = 250, alternative = "greater")
t.test(chickwts$weight, mu = 250, alternative = "less")
```

Note that hypothesis testing is concerned with finding evidence in support of the null hypothesis  $H_0$  - the default situation - rather than evidence in favour of the alternative hypothesis.

## One Sample t-test

For the chicken weights data an appropriate formal analyses is to use a **one-sample t-test**, why this test is appropriate will be discussed later. This analysis involves calculating a simple summary statistic - called a *t*-statistic - which we do entirely from the observed data.

$$T_{obs} = rac{ar{x} - \mu}{s / \sqrt{n}}$$

where  $\bar{x}$  is the sample mean, s the sample standard deviation and  $\mu$  is the population mean in the null hypothesis which we wish to test for. We then look up the value of  $T_{obs}$  in a set of statistical tables/computer to see what the answer is to our research question.

## Important concept - sampling

Why is 
$$T_{obs} = \frac{\bar{x} - \mu}{s / \sqrt{(n)}}$$
 called a *t*-statistic?

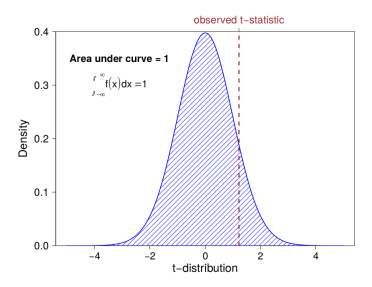
If another sample of 71 chickens from the same population were weighed then the values for  $\bar{x}$  and s would be different, and hence the value for  $T_{obs}$ . If this was repeated many times and a histogram/Q-Q/P-P plot produced of the values for  $T_{obs}$  then this would follow the shape of a known distribution - **student-***t* **probability distribution**. It is this piece of mathematics - knowing what the sampling distribution of  $T_{obs}$  is - which allows us to infer information about the population of chickens from which our original 71 chickens were sampled - without actually having to collect lots and lots of other samples of chickens! Mathematical theory is used to fill this data gap.

#### chickwts: t-test

$$T_{obs} = \frac{261.31 - 250}{78.07/\sqrt{71}} = 1.22$$

Put the values for the sample mean and standard deviation into the t-statistic formula along with the  $\mu=250$ . We now look up the value of this in a t-distribution reference table. All this calculation will be done for you in R but it is important to understand the general process as this is the same for hypothesis testing in other more complex analyses.

## One Sample, one-sided, t-test



## Important concept - p-values

- The result of a hypothesis test is usually communicated in the form of a p-value
- The interpretation of a p-value is of crucial importance it is the probability that the test statistic takes values at least as extreme as that observed assuming that H<sub>0</sub> is true
- Exactly what at least as extreme as refers to depends on the alternative hypothesis H<sub>A</sub>.
- This may sound rather abstract but it is usually obvious in practice

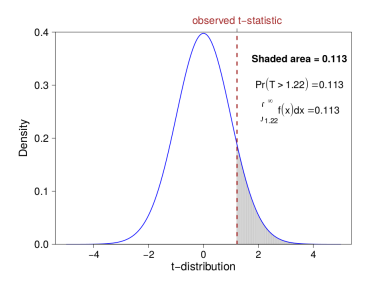
After six weeks will the mean weight of a chicken be at least 250 grams?

 $H_0: \mu \leq 250$ 

 $H_A: \mu > 250$ 

The alternative hypothesis is  $\mu >$  250 so in this test **at least as extreme as** in the definition of a p-value is the probability of observing a t-statistic which is > 1.22 **assuming that**  $H_0$  **is true** - this is why 250 was used for  $\mu$  when calculating  $T_{obs}$ .

## One Sample, one-sided, t-test



#### **Research Question**

The purpose of this hypothesis test analysis is to answer a very specific scientific question:

## After six weeks will the mean weight of a chicken be more than 250 grams?

So what is our answer?

The *p*-value for this hypothesis test is 0.113. Based on this value we can either **reject**  $H_0$  and conclude that the mean weight of chickens in the population is likely to be greater than 250 grams or else we can **accept**  $H_0$  where the mean chicken weight is less than 250 grams.

## Research Question - be pragmatic with p-values

By convention a p-value of less than 0.05 is considered to provide **reasonable evidence** for rejecting  $H_0$ . A p-value of between 0.05 and 0.1 might be considered as **weak evidence** against  $H_0$ . Values of less that 0.01 are generally considered as **very strong evidence** for rejecting  $H_0$ . It is **always** best to provide a p-value in any analyses to let the reviewer/client see the strength of evidence rather than simply claiming statistically significant findings!

## **Communicating Results of Hypothesis Tests**

Transparency is essential - the devil can be in the detail - which at the very least should comprise:

- i. what hypothesis was being tested be clear and precise
- ii. what statistical test was used
- iii. what the p-value is
- iv. what the treatment effect is (more later).

This is particularly crucial if the analyses are to be given to someone *else* to then make a judgment on the scientific significance.

## **Two-sided Tests: One Population**

## After six weeks will the mean weight of a chicken be equal to 250 grams?

This is now a two sided hypothesis test:

 $H_0: \mu = 250$ 

 $H_A: \mu \neq 250$ 

This time the hypothesis test is asking how much evidence is there in our sample data to conclude that in the population of all chickens the mean weight is not equal to 250 grams.

## **Two-sided Tests: One Population**

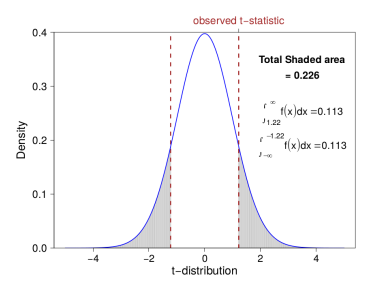


```
# t.test(chickwts$weight, mu = 250, alternative = "two.sided")
t.test(chickwts$weight, mu = 250)
##
##
   One Sample t-test
##
## data: chickwts$weight
## t = 1.2206, df = 70, p-value = 0.2263
## alternative hypothesis: true mean is not equal to 250
## 95 percent confidence interval:
## 242.8301 279.7896
## sample estimates:
## mean of x
## 261.3099
```

#### **Two-sided Tests**

A two-sided test is similar to a one-sided test - the key difference is in what is now defined as **at least as extreme** in the definition of the p-value. This time the alternative hypothesis refers to observing a value of **either**  $\bar{x} > 250$  or  $\bar{x} < -250$  **assuming that**  $H_0$  **is true**, which using the t-test approach is equivalent to the probability of observing  $T_{obs} > 1.22$  or  $T_{obs} < -1.22$  which we can again look up in reference tables.

## One Sample, two-sided, t-test



#### **Two-sided Tests**

- The two-sided t-test has a p-value which is exactly double the single sided test.
- Think! Intuitively the p-value should be less for a single sided test as the research question you are asking is much narrower e.g. greater than 250 grams, as opposed to whether the mean chicken weight might be either less than 250 grams or greater than 250 grams.
  - $\rightarrow$  You are using the same amount of information (71 observations) to answer a narrower research question and so all else being equal you should expect a "more powerful" analyses (e.g. a lower *p*-value all else being equal)

## **Exercise 11**



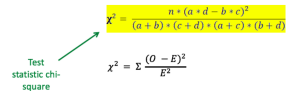
## **Chi-square Test**

There are two very commonly used statistical tests for testing dependence between two categorical variables: Chi-squared test & Fisher's exact test.

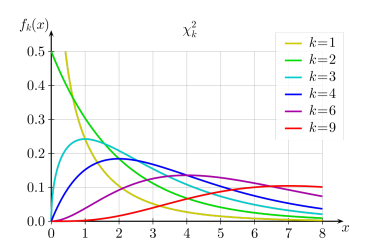
## To test independence of rows and columns

	Disease		
Risk Factor	+	-	Total
+	a	b	a+b
-	С	d	c+d
Total	a+c	b+d	n = a+b+c+d

• Assumptions: a, b, c, d must have at least 5 observations!



## **The Chi-square Distribution**



## **Exact Fisher Test: Permutation Test**

	Success	Failure	Total
Therapy	7	2	9
New Therapy	2	8	10
Total	9	10	19

7 2 8	P = 9!*10!*9!*10! / 19!*7!*2!*2!*8! = 0.01754
8 1 1 9	P = 9!*10!*9!*10! / 19!*8!*1!*1!*9! = 0.00097
9 0 0 10	P = 9!*10!*9!*10! / 19!*9!*0!*0!*10! = 0.00001

For a one-sided test: p = 0.01754+0.00097+0.00001 = 0.01852

## **Exercise 12**



## Simulating in R



#### Continuous variable

```
n <- 100
rnorm(n, mean = 0, sd = 1)  # Normal distribution</pre>
```

## Binary variable

```
rbinom(n, size = 1, prob = 0.4) # Binomial distribution
```

#### Count variable

```
rpois(n, lambda = 7)  # Poisson distribution
```

## Other options

```
seq(from = 0, to = 100, by = 1)  # ID
sample(3:30, size = n, replace = TRUE)  # herd size
c(rep(c(1, 2, 3), times = 33), 1)
c(rep(c(1, 2, 3), each = 33), 1)
```

•

## **Side remark: Conditional Inference Trees (1/2)**



 ctree(...) is a non-parametric class of regression trees embedding tree structured regression models into a well defined theory of conditional inference procedures.

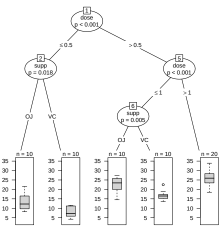
## Recursive partitioning algorithm:

- partition the observations by univariate splits in a recursive way
- fit a constant model in each cell of the resulting partition
- $\bullet \to \text{uses}$  an information measure of node impurity to select the variables showing the best split

## **Side remark: Conditional Inference Trees (2/2)**



```
library("partykit")
library("party")
data("ToothGrowth")
my.tooth.tree <- ctree(len ~ ., data = ToothGrowth)
par(mfrow=c(1,1))
plot(my.tooth.tree, tp_args = list(id = FALSE))</pre>
```



## Plotting in R



## **Plotting in R**



- Continuous data
  - Histogram
  - Boxplot
- Nominal / Ordinal data
  - Barplot
  - Mosaicplot
  - Scatterplots

## Exercise 13A and 13B



## **Exercise 14**



#### Overview: ANOVA and linear models

- Introduction to ANOVA
- How to run an ANOVA in R
- Checking model assumptions in R
- Multiple comparisons options in R
- ANOVA as a special case of a linear model
- The simple linear regression model
- The multiple linear regression model
- Model selection: R<sup>2</sup> and AIC
- Two-way Interactions in R
- Confounding

#### **Hypothesis Testing - One Way ANOVA**

We have seen how to perform hypothesis tests when comparing two populations using the two sample t-test. In many analyses we may have multiple populations - for example suppose we have a treatment which has a number of different levels high/medium/low/placebo, or equivalently a number of different treatments. What then is the hypothesis we wish to test?

#### Is there a difference in the effect of the treatment?

 $H_0$ :  $\mu_1 = \mu_2 = \cdots = \mu_k$ 

 ${\it H}_{\it A}$  : at least one pair of  $\mu_1,\ldots,\mu_k$  are different

where  $\mu_1, \ldots, \mu_k$  denote the mean effect of treatment levels 1 through k.

#### **One Way ANOVA**

Analysis of variance, **ANOVA**, to analyze differences between group means. The observed variance in the outcome variable is partitioned into components attributable to different sources of variation.

### ANOVA estimates three sample variances (sum of squares)

- a total variance based on all observation deviations from the grand mean
- a variance based on the group mean deviations from the grand mean
- an (error) variance based on all the observation deviations from their group mean

$$\sum_{i} \sum_{j} (x_{ij} - \bar{x})^2 = \sum_{i} n_i (\bar{x}_i - \bar{x})^2 + \sum_{i} \sum_{j} (x_{ij} - \bar{x}_i)^2$$

Variance between groups

Variance within groups

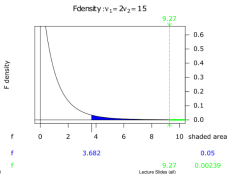
 $\bar{x}$ = grand mean  $\bar{x}_i$ = group mean

#### F-test / F-distribution

An **F-test**, a statistical test, in which the test statistic has an F-distribution under the null hypothesis is used to assess statistical significance in an ANOVA.

degrees of freedom

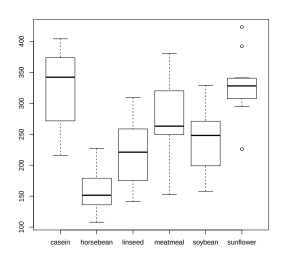
$$\mathsf{F} = \frac{variance\ between\ groups}{variance\ within\ groups}$$



#### ANOVA in R with chickwts



data(chickwts)
boxplot(chickwts\$weight ~ chickwts\$feed)



#### ANOVA in R with chickwts



```
# aov.mod <- aov(chickwts$weight ~ chickwts$feed)
aov.mod <- aov(weight ~ feed, data = chickwts)</pre>
# What objects can we extract from a anova model?
objects(aov.mod)
## [1] "assign"
                     "call"
                                     "coefficients" "contrasts"
## [5] "df.residual" "effects" "fitted.values" "model"
## [9] "qr"
                      "rank"
                                     "residuals" "terms"
## [13] "xlevels"
#
summary(aov.mod)
##
             Df Sum Sq Mean Sq F value Pr(>F)
## feed 5 231129 46226 15.37 5.94e-10 ***
## Residuals 65 195556 3009
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

#### **Overview: Checking model assumptions**

- mean(residuals) = 0
- Residuals are normally distributed (qqnorm, qqplot)
- Variance homoscedasticity (Bartlett & Levene's Test)
- Cook's distance: Influential data points
- Any pattern(s)?

### Checking model residuals: mean(residuals) = 0

# R

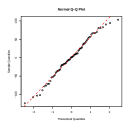
#### "Unexplained rest of the model"

```
# What are residuals?
chickwts$residuals <- residuals(aov.mod)
tapply(chickwts$weight, chickwts$feed, mean)
     casein horsebean linseed meatmeal sovbean sunflower
##
## 323.5833 160.2000 218.7500 276.9091 246.4286 328.9167
chickwts[c(1:3).]
    weight feed residuals
##
## 1 179 horsebean 18.8
## 2 160 horsebean -0.2
## 3 136 horsebean -24.2
# Save residuals to an objects and check mean of residuals
aov.mod.resid <- residuals(aov.mod)
mean(aov.mod.resid)
## [1] 7.573045e-16
```

# Checking model residuals: Residuals normally distributed "Unexplained rest of the model"



```
par(mfrow=c(1,1))
qqnorm(aov.mod.resid)
qqline(aov.mod.resid, col = "red", lwd = 3, lty = 2)
```



```
# Shapiro-Wilk test (dependent on sample size --> limited use)
shapiro.test(aov.mod.resid)

##
## Shapiro-Wilk normality test
##
## data: aov.mod.resid
## W = 0.98616, p-value = 0.6272
```

# Checking model residuals: Variance homoscedasticity (1/3) Hypothesis tests



```
# Bartlett Test
bartlett.test(chickwts$weight ~ chickwts$feed)

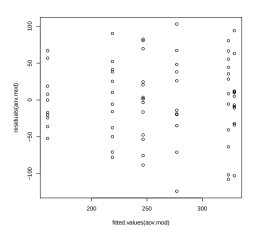
##
## Bartlett test of homogeneity of variances
##
## data: chickwts$weight by chickwts$feed
## Bartlett's K-squared = 3.2597, df = 5, p-value = 0.66

# Levene's Test
# library("Rcmdr")
# levene.test(chickwts$weight ~ chickwts$feed)
```

# Checking model residuals: Variance homoscedasticity (2/3) Graphical interpretation is better!



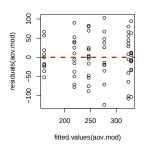
```
# Plot fitted against residual values
plot(fitted.values(aov.mod), residuals(aov.mod))
```

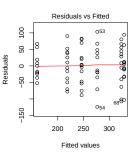


# Checking model residuals: Variance homoscedasticity (3/3) Graphical interpretation is better!



```
# Plot fitted against residual values
par(mfrow=c(1,2), pty="s", mar = c(10, 4, 1, 2))
plot(fitted.values(aov.mod), residuals(aov.mod))
abline(h = 0, col = "red", lwd = 3, lty = 2)
plot(aov.mod, which=1)
```

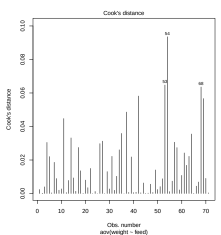




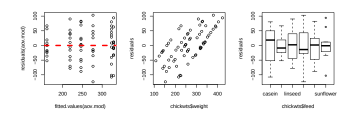
### Checking model residuals: Cook's distance



```
# Plot fitted against residual values
# Cut-off at 3
par(mfrow=c(1,1), pty="s", mar = c(5, 4, 4, 2))
plot(aov.mod, which=4)
```



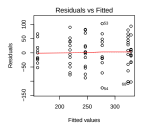
#### **Checking for potential patterns**

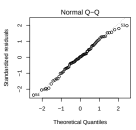


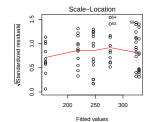
#### Plot anova objects

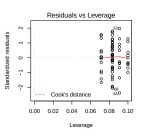
par(mfrow=c(2, 2))
plot(aov.mod)











### Multiple comparisons options in R



So far, we know there is difference between the feed types. However, we do not yet know which feed type differ. In principal, we could do multiple t-tests, BUT ... we would use our data several times. Classically, we choose an  $\alpha$ -level of 5 %, in cases of multiple comparisons we are facing the familywise error rate:

$$FWE \leq 1 - (1 - \alpha)^n,$$

where  $\alpha = 5$  % and n = number of comparisons.

```
alpha = 0.05

1 - (1 - alpha)^1  # n = 1  # [1] 0.05

1 - (1 - alpha)^5  # n = 5  # [1] 0.2262191

1 - (1 - alpha)^10  # n = 10  # [1] 0.4012631
```

#### Multiple comparisons options in R



Hence, we are better of using one of the following procedures to adjust for multiple comparisons:

- **Bonferroni**: p-value correction by testing each individual hypothesis at a significance level of  $\frac{\alpha}{m}$ , where  $\alpha$  is the desired overall  $\alpha$  level and m is the number of hypotheses.
- **Dunnett**: Multiple comparisons of each group to a reference.
- Tukey Honest Significant Differences (Homogeneous subgroups): Multiple comparisons of all possible combinations.

• ...

### Multiple comparisons options in R: Bonferroni



```
aov.mod <- aov(weight ~ feed, data = chickwts)
pairwise.t.test(chickwts$weight, chickwts$feed, p.adj = "none")
##
   Pairwise comparisons using t tests with pooled SD
##
## data: chickwts$weight and chickwts$feed
##
            casein horsebean linseed meatmeal sovbean
##
## horsebean 2.1e-09 -
## linseed 1.5e-05 0.01522 -
## meatmeal 0.04557 7.5e-06 0.01348 -
## soybean 0.00067 0.00032 0.20414 0.17255 -
## sunflower 0.81249 8.2e-10 6.2e-06 0.02644 0.00030
##
## P value adjustment method: none
pairwise.t.test(chickwts$weight, chickwts$feed, p.adj = "bonferroni")
##
   Pairwise comparisons using t tests with pooled SD
##
## data: chickwts$weight and chickwts$feed
##
##
            casein horsebean linseed meatmeal sovbean
## horsebean 3.1e-08 -
## linseed 0.00022 0.22833 -
## meatmeal 0.68350 0.00011 0.20218 -
## soybean 0.00998 0.00487 1.00000 1.00000 -
## sunflower 1.00000 1.2e-08 9.3e-05 0.39653 0.00447
##
## P value adjustment method: bonferroni
```

#### Multiple comparisons options in R: Dunnett



```
library("multcomp")
# compares always to baseline levels (here: casein) --> saves degrees of freedom
dunnett <- glht(aov.mod, linfct = mcp(feed = "Dunnett"))</pre>
summary(dunnett)
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Multiple Comparisons of Means: Dunnett Contrasts
##
##
## Fit: aov(formula = weight ~ feed, data = chickwts)
##
## Linear Hypotheses:
                         Estimate Std. Error t value Pr(>|t|)
##
## horsebean - casein == 0 -163.383
                                      23.485 -6.957 < 0.001 ***
## linseed - casein == 0 -104.833 22.393 -4.682 < 0.001 ***
## meatmeal - casein == 0 -46.674 22.896 -2.039 0.16715
## sovbean - casein == 0 -77.155 21.578 -3.576 0.00314 **
## sunflower - casein == 0.5.333 22.393 0.238 0.99945
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
```

# Multiple comparisons options in R: Tukey Tukey Honest Significant Differences

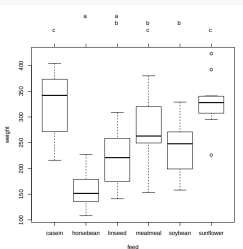


```
library("multcomp")
# compares all factor levels
tukey <- glht(aov.mod, linfct = mcp(feed = "Tukey"))</pre>
summarv(tukev)
##
##
    Simultaneous Tests for General Linear Hypotheses
##
## Multiple Comparisons of Means: Tukey Contrasts
##
##
## Fit: aov(formula = weight ~ feed, data = chickwts)
##
## Linear Hypotheses:
##
                             Estimate Std. Error t value Pr(>|t|)
## horsebean - casein == 0
                             -163.383
                                          23.485 -6.957 < 0.001 ***
                             -104.833
## linseed - casein == 0
                                          22.393 -4.682 < 0.001 ***
## meatmeal - casein == 0
                            -46.674
                                          22.896 -2.039 0.33209
## soybean - casein == 0
                              -77.155
                                          21.578 -3.576 0.00845 **
## sunflower - casein == 0
                              5.333
                                          22.393
                                                   0.238 0.99989
## linseed - horsebean == 0
                               58.550
                                          23.485
                                                   2.493 0.14132
                                          23.966
## meatmeal - horsebean == 0
                              116.709
                                                   4.870 < 0.001 ***
## soybean - horsebean == 0
                              86.229
                                          22.710
                                                   3.797 0.00423 **
## sunflower - horsebean == 0
                              168.717
                                          23.485
                                                   7.184 < 0.001 ***
## meatmeal - linseed == 0
                               58.159
                                          22.896
                                                   2.540 0.12749
## soybean - linseed == 0
                              27.679
                                          21.578
                                                   1.283 0.79294
## sunflower - linseed == 0
                              110.167
                                          22.393
                                                   4.920 < 0.001 ***
## soybean - meatmeal == 0
                              -30.481
                                          22.100 -1.379 0.73879
## sunflower - meatmeal == 0
                              52.008
                                                   2.271 0.22030
                                          22.896
## sunflower - soybean == 0
                              82.488
                                          21.578
                                                   3.823 0.00389 **
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
statia Hakirikadı, TerkmeriDotoch & Nülriel និសា reported -- single-step method) Lecture Slides (all)
```

# Multiple comparisons options in R: Tukey Tukey Honest Significant Differences with homogeneous subgroups



```
# summary(tukey) # standard display
tukey.cld <- cld(tukey) # letter-based display
# the cld(...) function sets up a compact letter display of all pair-wise comparisons
par(mfrow=c(1,1), mar=c(5, 4, 5, 2))
plot(tukey.cld)</pre>
```



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#### **Exercise 15**



### Introduction: ANOVA as a special case of a linear model Linear Modelling

So far we have considered how to determine whether statistically significant differences exist between different "feed" groups (factors).

The explanatory variable has been categorical. We have not yet considered continuous explanatory variables.

ANOVA and linear regression are both a special cases of **linear** models.

#### Simple linear regression (1/3)

A **simple linear regression** fits a straight line through a set of data points. This straight line is fitted in a way that makes the sum of the squared residuals (the vertical distances between each data point and the fitted line) as small as possible.

#### Simple linear regression (2/3)

In a simple linear regression model for n data points  $(x_i, y_i)$ , i = 1, ...n the following equation is used:

$$y_i = \alpha + \beta x_i + \epsilon$$
,

#### where

- α: intercept or constant
- $\beta$ : slope, regression coefficient (effect size)
- $\epsilon$ : error, residuals

#### Simple linear regression (3/3)

The goal is to find the equation of the straight line

$$f(x) = y = \alpha + \beta x$$
,

which would provide a "best" fit for the data points (least square approach).

#### **Assumptions of a linear regression**

A **simple linear regression** is based on several assumptions which should be checked carefully.

- linearity of the relationship between the explanatory (independent) and the outcome (dependent) variable
- normality of the residuals
- independence
- constant variance (homoscedasticity)

### Linear regression model in R

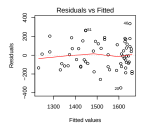


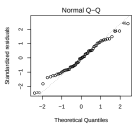
```
data(water)
# mod.hard <- lm(mortality ~ hardness, data = water)</pre>
mod.hard <- lm(water$mortality ~ water$hardness)</pre>
summary(mod.hard)
##
## Call:
## lm(formula = water$mortality ~ water$hardness)
##
## Residuals:
## Min 1Q Median 3Q Max
## -348.61 -114.52 -7.09 111.52 336.45
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1676.3556 29.2981 57.217 < 2e-16 ***
## water$hardness -3.2261 0.4847 -6.656 1.03e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 143 on 59 degrees of freedom
## Multiple R-squared: 0.4288, Adjusted R-squared: 0.4191
## F-statistic: 44.3 on 1 and 59 DF, p-value: 1.033e-08
```

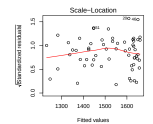
### **Checking linear model assumptions**

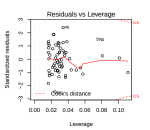
R

par(mfrow=c(2,2))
plot(mod.hard)









#### Linear model vs. t.test(...) in R



```
mod.loc <- lm(mortality ~ location, data = water)</pre>
coef(mod.loc)
##
     (Intercept) locationSouth
##
       1633.6000 -256.7923
t.test(water$mortality ~ water$location)
##
##
    Welch Two Sample t-test
##
## data: water$mortality by water$location
## t = 7.1427, df = 53.29, p-value = 2.584e-09
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 184.6919 328.8928
## sample estimates:
## mean in group North mean in group South
##
              1633,600
                           1376.808
```

# The multiple linear model An extension of the simple linear model (1/2)

If several explanatory variables are of interest, instead of performing multiple simple regressions, a **multiple linear regression** or **multivariable** approach is appropriate.

- the same assumptions as for simple linear regressions should be checked
- collinearity might be an issue (see vif(...) function from package usdm)
- model comparison (AIC) (...discussed later)

# The multiple linear model An extension of the simple linear model (2/2)

Simple linear regression model:

$$y_i = \alpha + \beta x_i + \epsilon$$

Multiple linear regression model:

$$y_i = \alpha + \beta_1 x \mathbf{1}_i + \beta_2 x \mathbf{2}_i + \dots + \epsilon,$$
  

$$y_i = \alpha + \beta_1 x \mathbf{1}_i * \beta_2 x \mathbf{2}_i + \dots + \epsilon,$$
(1)

#### where

- *α*: intercept or constant
- β<sub>1</sub>: slope1, regression coefficient (effect size)
- β<sub>2</sub>: slope2, regression coefficient (effect size)
- $\beta_1 * \beta_2$ : interaction effect between  $x_1$  and  $x_2$
- $\epsilon$ : error, residuals

## The multiple linear model Interpretation of model coefficients

- α: intercept or constant
- β<sub>1</sub>: slope1, regression coefficient (effect size)
- β<sub>2</sub>: slope2, regression coefficient (effect size)
- $\beta_1 * \beta_2$ : interaction effect between  $x_1$  and  $x_2$
- $\epsilon$ : error, residuals
- → β<sub>1</sub> describes the number of units of a change in the outcome variable y as x<sub>1</sub> changes by one unit, x<sub>2</sub> being held constant.
- → β<sub>2</sub> describes the number of units of a change in the
   outcome variable y as x<sub>2</sub> changes by one unit, x<sub>1</sub> being held
   constant.

#### The multiple linear model in R



```
mod.hard.loc <- lm(mortality ~ hardness + location, data = water)
summary(mod.hard.loc)
##
## Call:
## lm(formula = mortality ~ hardness + location, data = water)
##
## Residuals:
## Min 10 Median 30
                                        Max
## -222.959 -77.281 7.143 90.751 307.836
##
## Coefficients:
##
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1695.4371 25.3285 66.938 < 2e-16 ***
## hardness -2.0341 0.4829 -4.212 8.93e-05 ***
## locationSouth -176.7108 36.8913 -4.790 1.19e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 122.1 on 58 degrees of freedom
## Multiple R-squared: 0.5907, Adjusted R-squared: 0.5766
## F-statistic: 41.86 on 2 and 58 DF, p-value: 5.601e-12
```

#### **Model selection**



- Multiple  $R^2$ : the larger, the better
- Akaike criterion (AIC): the smaller, the better

#### **Nested models**



Two models are nested if one of them is a particular case of the other one: the simpler model can be obtained by setting some coefficients of the more complex model to particular values.

#### Among the 3 above models ...

- which ones are nested?
- which ones are not nested?

# Multiple R<sup>2</sup>



 $R^2$  is a measure of fit quality:

$$R^2 = rac{ ext{SSRegression}}{ ext{SSTotal}}$$
 $R^2 = 1 - rac{ ext{SSError}}{ ext{SSTotal}}$ 

#### CAREFUL:

- The SSError always decreases as more predictors are added to the model.
- R<sup>2</sup> always increases and can be artificially large.

#### Akaike criterion (AIC)

- Model fit (R<sup>2</sup>) always improves with model complexity. We would like to strike a good balance between model fit and model simplicity.
- AIC combines a measure of model fit with a measure of model complexity: The smaller, the better.

For a given data set and a given model:

$$AIC = -2 \cdot \log(L) + 2p$$

L stands for the likelihood. p stands for the number of parameters in the models (penalizes complex models).

#### Model selection strategy for AIC

- Consider a number of candidate models. (They need not be nested.)
- Calculate their AIC.
- Choose the model(s) with the smallest AIC.
- ightarrow **CAREFUL**: The absolute value of AIC is meaningless. The relative AIC values, between models, is meaningful.

#### Model selection with AIC in R

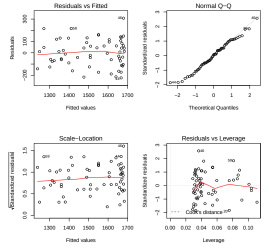


```
mod1 <- lm(mortality ~ hardness, data = water)</pre>
mod2 <- lm(mortality ~ location, data = water)</pre>
mod3 <- lm(mortality ~ hardness + location, data = water)
AIC(mod1, mod2, mod3)
## df AIC
## mod1 3 782.5692
## mod2 3 778.5186
## mod3 4 764.2366
round(AIC(mod1, mod2, mod3), 2)
## df AIC
## mod1 3 782.57
## mod2 3 778.52
## mod3 4 764.24
```

#### mod.hard.loc is the best!



```
\label{eq:mod3} $$ $$ \mbox{mod3} <-\mbox{lm(mortality $\tilde{\mbox{$^{\circ}$}}$ hardness $$+$ location, data = water)} $$ $$ \mbox{par(mfrow=c(2,2))} $$ $$ \mbox{plot(mod3)} $$
```



#### **Exercise 16**



## Two-way Interactions in R (1/3)



#### There are three different interactions:

- Interaction between two categorical variables
- Interaction between one continuous and one categorical variables
- Interaction between two continuous variables

## Two-way Interactions in R (2/3)



# Model specification in R: Be aware, an interaction is never tested without its corresponding main effects included in the model.

```
# Interaction between two categorical variables
# mod.dose.supp.int <- lm(len ~ dose.fac + supp + dose.fac:supp,
# data = ToothGrowth)
mod.dose.supp.int <- lm(len ~ dose.fac * supp, data = ToothGrowth)
# summary(mod.dose.supp.int)</pre>
```

$$\begin{split} \textit{y} \sim \beta_{\text{baseline}((\text{dose==low}) \& (\text{supp==OJ}))} + \beta_{\text{dose==med}} + \\ \beta_{\text{dose==high}} + \beta_{\text{supp==VC}} \\ + \beta_{(\text{dose==med}) \& (\text{supp==VC})} \\ + \beta_{(\text{dose==high}) \& (\text{supp==VC})} \end{split}$$

#### Two-way Interactions in R (3/3)



```
# Interaction between two categorical variables
# mod.dose.supp.int <- lm(len ~ dose.fac + supp + dose.fac:supp.
# data = ToothGrowth)
mod.dose.supp.int <- lm(len ~ dose.fac * supp, data = ToothGrowth)
summarv(mod.dose.supp.int)
##
## Call:
## lm(formula = len ~ dose.fac * supp. data = ToothGrowth)
##
## Residuals:
##
     Min 1Q Median 3Q
                               Max
## -8.20 -2.72 -0.27 2.65 8.27
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                      13.230 1.148 11.521 3.60e-16 ***
## dose.facmed
                       9.470
                                  1.624 5.831 3.18e-07 ***
                      12.830 1.624 7.900 1.43e-10 ***
## dose.fachigh
                      -5.250 1.624 -3.233 0.00209 **
## suppVC
## dose.facmed:suppVC -0.680
                                  2.297 -0.296 0.76831
## dose.fachigh:suppVC
                      5.330
                                  2 297 2 321 0 02411 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 3.631 on 54 degrees of freedom
## Multiple R-squared: 0.7937, Adjusted R-squared: 0.7746
## F-statistic: 41.56 on 5 and 54 DF, p-value: < 2.2e-16
```

## Two-way Interactions in R: Interpretation of coefficients (1/2)



```
## (Intercept) dose.facmed dose.fachigh
## 13.23 9.47 12.83
## suppVC dose.facmed:suppVC dose.fachigh:suppVC
## -5.25 -0.68 5.33
```

$$y \sim eta_{
m baseline((dose==low)\ \&\ (supp==OJ))} + eta_{
m dose==med} + \ eta_{
m dose==high} + eta_{
m supp==VC} \ + eta_{
m (dose==med)\ \&\ (supp==VC)} \ + eta_{
m (dose==high)\ \&\ (supp==VC)}$$

#### Two-way Interactions in R: Interpretation of coefficients (2/2)



```
coef(mod.dose.supp.int)

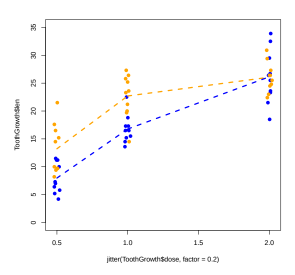
## (Intercept) dose.facmed dose.fachigh
## 13.23 9.47 12.83

## suppVC dose.facmeds:uppVC dose.fachigh:suppVC
## -5.25 -0.68 5.33
```

- → dose.facmed:suppVC change in the slope between the low and med dose.fac group under the supplement type (supp) VC in comparison to the intercept. In other words, by changing the dose from low to med within the supplement group VC, the slope decreases by approx. -0.68.
- dose.fachigh:suppVC change in the slope between the low and high dose.fac group under the supplement type (supp) VC in comparison to the intercept. In other words, by changing the dose from low to high within the supplement group VC, the slope increases by approx. +5.33.

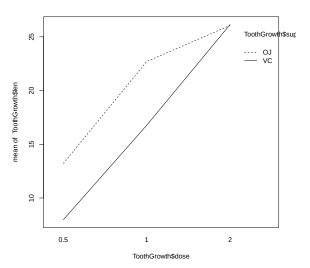
#### **Interactions**





#### **Interactions**





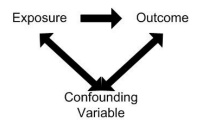
#### **Exercise 17**



#### Confounding



- Confounding variable is associated with the exposure and the outcome
- Confounding variable is not part of the causal path between exposure and the outcome



#### Some things to remember

- Not adjusting / controlling for a confounding variable may lead to biased results. It is good practice to present as well the crude (not adjusted ORs) and the adjusted ones. Adjustment is typically done if difference > 10%.
- Check also for interaction (= effect modification) e. g. using logistic regression.
- Do not adjust for a variable C if it is a common effect of E and D (collider) or if it is in the causal pathway of E and D.

#### **Exercise 18**



#### Other topics

So far, we have covered some of the basic **tools** necessary for helping you to start to analyse your own study data.

Many things have been mentioned others have not been included some are worth mentioning as they are particularly common in some areas of veterinary research.

#### Normally distributed response variables

So far, we have only considered analyses where the response variable is a **continuous** measurement.

We now have a brief look at two other common types of **response** variables.

The methods we use are largely similar to what we have looked at previously.

### Data types in R



#### numeric & integers

```
ToothGrowth$len[1:6]
## [1] 4.2 11.5 7.3 5.8 6.4 10.0
bacteria$week[1:6]
## [1] 0 2 4 11 0 2
```

#### unordered factor with 2 levels

```
lung$sex[1:6]
## [1] female male female female male female
## Levels: female male
```

#### (un/ordered) factor (with more than 2 levels)

```
chickwts$feed[1:6]
## [1] horsebean horsebean horsebean horsebean horsebean
## Levels: casein horsebean linseed meatmeal soybean sunflower
```

#### **Generalised Linear Modelling**

- random component:  $\mathbf{E}(\mathbf{Y}) = \mu$
- systematic component: covariates:  $\mathbf{x_1}, \mathbf{x_2}, \dots, \mathbf{x_p}$  produce a linear predictor  $\eta = \sum_1^p \mathbf{x_j} \beta_j$
- link between random and systematic components:  $\mu = \eta$ , e.g.
  - Logistic Regression:  $\eta_i = g(\mu_i), g(.)$  is logit function for binomial data
  - Poisson Regression:  $\eta_i = g(\mu_i), g(.)$  is exponential function for poisson data

# Logistic Regression: unordered factor with 2 levels as response

E. g. a logistic regression with one covariate (sex) fitting a model to  $\mathbb{P}(Y = TRUE) = \pi$ , e. g. presence of respiratory symptoms (experienced by the child over the previous 12 months), gives

$$\log\left(\frac{\pi}{1-\pi}\right) = \beta_0 + \beta_1 \mathbf{x}_1$$

# Logistic Regression: unordered factor with 2 levels as response

```
logreg.mod <- glm(respsymptoms ~ sex.
                 family = binomial, # description of the link function
                 data = lung)
summary(logreg.mod)
##
## Call:
## glm(formula = respsymptoms ~ sex, family = binomial, data = lung)
## Deviance Residuals:
      Min
               1Q Median 3Q
                                        Max
## -0.7440 -0.7440 -0.6915 -0.6915 1.7597
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.1429 0.1276 -8.957 <2e-16 ***
## sexmale -0.1663 0.1901 -0.875 0.382
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 682.85 on 635 degrees of freedom
##
## Residual deviance: 682.08 on 634 degrees of freedom
## AIC: 686.08
##
## Number of Fisher Scoring iterations: 4
```

# Logistic Regression: unordered factor with 2 levels as response Interpretation of model coefficients

```
summary(logreg.mod)
##
## Call:
## glm(formula = respsymptoms ~ sex, family = binomial, data = lung)
##
## Deviance Residuals:
      Min
##
                10 Median
                                         Max
## -0.7440 -0.7440 -0.6915 -0.6915 1.7597
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -1.1429 0.1276 -8.957 <2e-16 ***
## sexmale -0.1663 0.1901 -0.875 0.382
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 682.85 on 635 degrees of freedom
##
## Residual deviance: 682.08 on 634 degrees of freedom
## ATC: 686.08
##
## Number of Fisher Scoring iterations: 4
cbind(logodds = coef(logreg.mod), odds = exp(coef(logreg.mod)))
                logodds
                             odds
## (Intercept) -1.1428851 0.3188976
## seymale
              -0.1662919 0.8467990
```

#### Odds ratios versus log odds ratios

- The logistic regression model in R outputs the log odds ratios.
- exp(log odds ratio) = odds ratio
- An odds ratio (OR) is a measure of association between an exposure (here: sex) and an outcome (here: respsymptoms).
   The OR represents the odds that an outcome will occur given a particular exposure, compared to the odds of the outcome occurring in the absence of that exposure.

		Outcome		
		Yes	No	
Predictor	Yes	А	В	$OR = \frac{(A*D)}{(B*C)}$
	No	С	D	

# Logistic Regression: unordered factor with 2 levels as response Interpretation of model coefficients (cont'd)

The coefficient  $\beta_{\text{sex}}$  is the estimated amount by which the log odds of respsymptoms would in-/decrease if sex is male. The log odds of respsymptoms when sex is female is just above in the first row ( $\beta_{(\text{Intercept})}$ , reference level of sex).

#### Logistic Regression: probability of success as response

- A researcher is examining beetle mortality after 5 hours of exposure to carbon disulphide, at various levels of concentration of the gas.
- Beetles were exposed to gaseous carbon disulphide at various concentrations (in mg/L) for five hours (Bliss, 1935) and the number of beetles killed were noted.

## Logistic Regression: probability of success as response



```
logreg.mod <- glm(cbind(Num.Killed, Num.Surv) ~ Dose,</pre>
                 family = binomial, # description of the link function
                 data = beetle)
summarv(logreg.mod)
##
## Call:
## glm(formula = cbind(Num.Killed, Num.Surv) ~ Dose, family = binomial,
      data = beetle)
##
##
## Deviance Residuals:
##
      Min
                10 Median
                                  30
                                          Max
## -1.5941 -0.3944 0.8329 1.2592 1.5940
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -60.717 5.181 -11.72 <2e-16 ***
## Dose
               34 270
                            2 912 11 77 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 284.202 on 7 degrees of freedom
##
## Residual deviance: 11.232 on 6 degrees of freedom
## ATC: 41.43
##
## Number of Fisher Scoring iterations: 4
```

# Logistic Regression: probability of success as response Interpretation of model coefficients



```
summary(logreg.mod)
##
## Call:
## glm(formula = cbind(Num.Killed, Num.Surv) ~ Dose, family = binomial,
       data = beetle)
##
## Deviance Residuals:
##
       Min
                 1Q Median 3Q
                                            Max
## -1.5941 -0.3944 0.8329 1.2592 1.5940
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -60.717 5.181 -11.72 <2e-16 ***
               34.270 2.912 11.77 <2e-16 ***
## Dose
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 284.202 on 7 degrees of freedom
##
## Residual deviance: 11.232 on 6 degrees of freedom
## ATC: 41.43
##
## Number of Fisher Scoring iterations: 4
cbind(logodds = coef(logreg.mod), odds = exp(coef(logreg.mod)))
                 logodds
## (Intercept) -60.71745 4.273114e-27
## Dose 34.27033 7.645631e+14
Sonja Hartnack, Terence Odoch & Muriel Buri
                                                       Lecture Slides (all)
```

# Logistic Regression: probability of success as response Interpretation of model coefficients (cont'd)



The coefficient  $\beta_{\text{Dose}}$  is the estimated amount by which the log odds of being killed would in-/decrease if Dose would increase by one unit. The log odds of being killed when Dose is 0 is just above in the first row ( $\beta_{\text{(Intercept)}}$ ).

#### Some things to remember

- Using generalised linear models (glm) is very similar to the previous models we have seen for normal data (lm) - things like residuals need to be checked for randomness, although this is more difficult with counts. Also the residuals being normally distributed is not relevant to these models.
- As always explore the data first visually and with tables before doing any formal analyses.