PHCM9795 Foundations of Biostatistics

Pilot Notes for R

05 July, 2022

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Introduction

These notes provide an introduction to R and instructions on how to conduct the analyses introduced in Foundations of Biostatistics.

These notes are currently under development, with sections being added and revised as the course progresses.

This is the first year that R has been offered as an option. I am keen to receive feedback about the notes and your experience learning R. Please get in touch if anything is unclear, or you have any questions or suggestions.

Changelog

2022-07-05

[Added]

· Module 6: first release.

2022-06-20

[Added]

· Module 5: first release.

2022-06-14

[Changed]

- Section 2.12 corrected the pnorm(q, mean, sd, lower.tail=FALSE) documentation to state that the it is the probablity of obtaining **more than** q that is calculated.
- Section 3.1 recommendation to use t.test() to calculate a 95% confidence interval for a mean, and not the descriptives() function as descriptives() uses a z-value instead of a t-value.

2022-06-10

[Added]

• Section 2.10 - Added instructions on labelling groups using the cut() function

2022-06-09

[Added]

• Section 2.8 - Summarising a single column of data using the descriptives() function from jmv package.

2022-06-07

[Changed]

• Section 2.6: Use the <- operator instead of =

4 CONTENTS

2022-06-05

[Changed]

· Module 1: Typos

2022-05-30

[Changed]

• Module 1: Typo in R Preferences (Section 1.3.1)

[Added]

Section 1.12: Instructions to plot a histogram with relative frequencies (i.e. percents) instead
of frequencies

2022-05-27

[Changed]

· Module 1: Fixed bar-charts that were not plotted correctly

2022-05-27

[Added]

- Section 1.2.1: Added a note about using the "patched" version of R 4.2.0 for Windows
- Section 1.14: Instructions for creating two-way tables using the contTables() function in the jmv package

2022-05-23

[Added]

Section 1.9: Explicit instructions to install jmv and summarytools when working in Module 1

[Changed]

 Section 1.9: Changed location of pbc.dat from examples to activities folder for consistency

2022-05-19

Initial release

Module 1

Introduction to R and RStudio

Learning outcomes

By the end of this Module, you will be able to:

- · understand the difference between R and RStudio
- · navigate the RStudio interface
- · input and import data into R
- · use R to summarise data
- perform basic data transformations
- · understand the difference between saving R data and saving R output
- copy R output to a standard word processing package

Part 1: An introduction to R

"R is a language and environment for statistical computing and graphics." Link. It is an open-source programming language, used mainly for statistics (including biostatistics) and data science. The aim of these notes is to introduce the R language within the RStudio environment, and to introduce the commands and procedures that are directly relevant to this course. There is so much more to R than we can cover in these notes. Relevant information will be provided throughout the course, and we will provide further references that you can explore if you are interested.

1.1 R vs RStudio

At its heart, R is a programming language. When you install R on your computer, you are installing the language and its resources, as well as a very basic interface for using R. You can write and run R code using the basic R app, but it's not recommended.

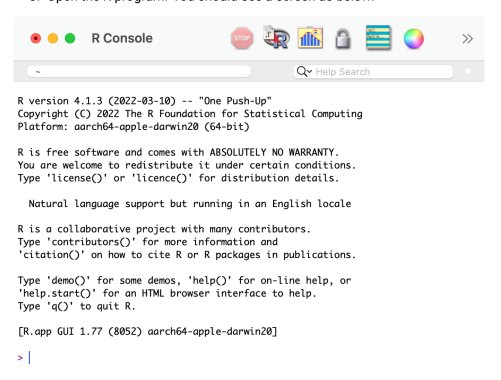
RStudio is an "Integrated Development Environment" that runs R while also providing useful tools to help you write code and analyse data. You can think of R as an engine which does the work, and RStudio as a car that uses the engine, but also provides useful tools like GPS navigation and reversing cameras that help you drive.

Note: even though we recommend that you use RStudio, you still need install R. **RStudio will not run without R installed.**

1.2 Installing R and RStudio

1.2.1 To install R on your computer

- 1. Download the R installer from:
 - a. for Windows: https://cran.r-project.org/bin/windows/base/
 - b. for MacOS: https://cran.r-project.org/bin/macosx/
 - Note for Windows users: as at May 27, 2022, R Version 4.2.0 has compatability issues with RStudio. You should download and install R from https://cran.r-project.org/bin/windows/base/rpatched.html
- 2. Install R by running the installer and following the installation instructions. The default settings are fine.
 - Note for macOS: if you are running macOS 10.8 or later, you will need to install an
 additional application called XQuartz, which is available at https://www.xquartz.org/.
 Download the latest installer (XQuartz-2.8.1.dmg as of April 2022), and install it in the
 usual way.
- 3. Open the R program. You should see a screen as below:



Near the bottom of the R screen, you will find the ">" symbol which represents the command line. If you type 1 + 2 into the command line and then hit enter you should get:

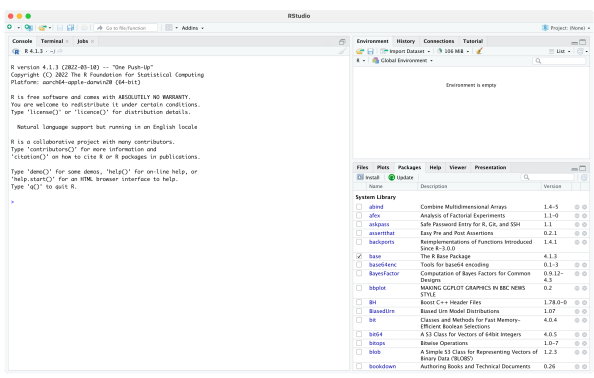
[1] 3

This is R performing your calculation, with the [1] indicating that the solution to 1 + 2 is a single number (the number 3).

At this point, close R - we will not interact with R like this in the future. You can close R by typing quit() at the command prompt, followed by the return key, or in the usual way of closing an application in your operating system. There is no need to save anything here if prompted.

1.2.2 To install RStudio on your computer

- 1. Make sure you have already installed R, and verified that it is working.
- Download the RStudio desktop installer at: https://www.rstudio.com/products/rstudio/download. Ensure that you select the RStudio Desktop (Free) installer in the first column.
- 3. Install RStudio by running the installer and following the installation instructions. The default settings are fine.
- 4. Open RStudio, which will appear as below:



Locate the command line symbol ">" at the bottom of the left-hand panel. Type 1 + 2 into the command line and hit enter, and you will see:

[1] 3

This confirms that RStudio is running correctly, and can use the R language to correctly calculate the sum between 1 and 2!

RStudio currently comprises three window panes, and we will discuss these later.

1.3 Recommended setup

I will provide a recommended setup for R and RStudio in this section. You are free to use alternative workflows and setup, but this setup works well in practice.

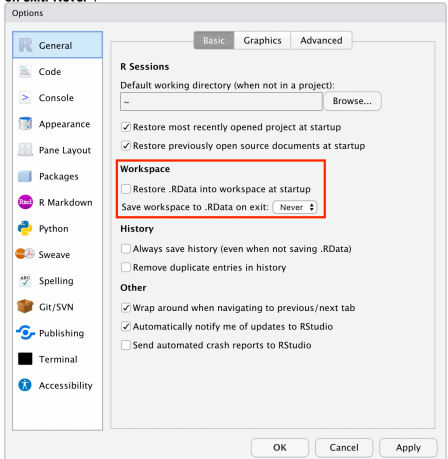
1.3.1 RStudio preferences

By default, RStudio will retain data, scripts and other objects when you quit your RStudio session. Relying on this can cause headaches, so I recommend that you set up RStudio so that it does not preserve your workspace between sessions. Open the RStudio options:

Mac: RStudio > Preferences

• Windows: Tools > Options

and deselect "Restore .RData into workplace at startup", and choose: "Save workspace to .RData on exit: Never".

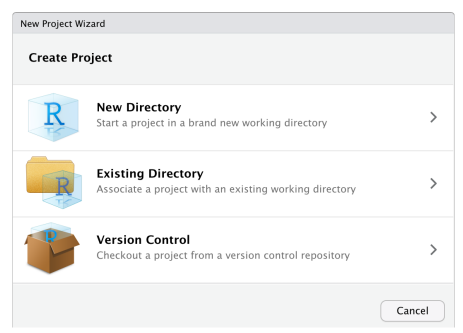


1.3.2 Set up a project

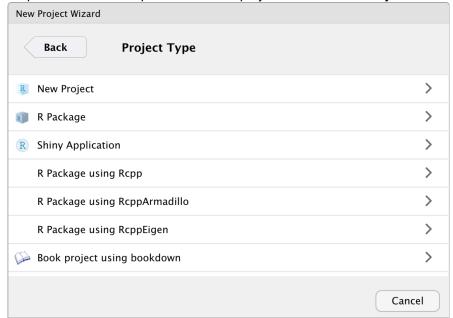
A project in RStudio is a folder that RStudio recognises as a place to store R scripts, data files, figures that are common to an analysis project. Setting up a folder allows much more simple navigation and specification of data files and output. More detail can be found in Chapter 8 of the excellent text: R for Data Science. Using projects is not necessary, but I recommend working with projects from day one.

We will create a project called **PHCM9795** to store all the data you will use and scripts that you will write in this course. First, think about where you want to store your project folder: this could be somewhere in your *Documents* folder.

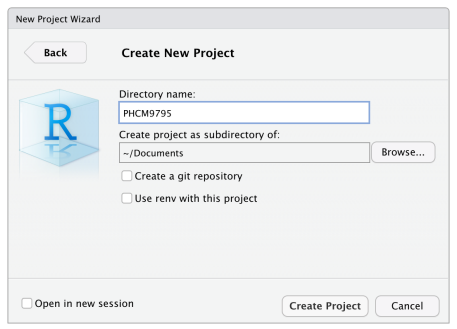
Step 1: Choose File > New Project... in RStudio to open the Create Project dialog box:



Step 2: Click the first option to create a project in a New directory

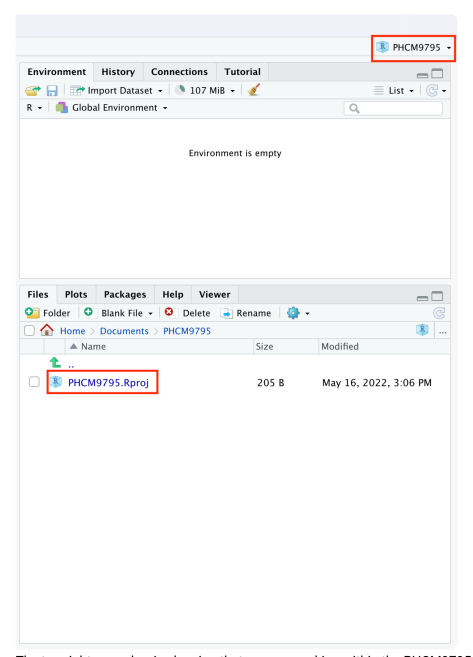


Step 3: Click the first option: **New Project**. Give the project a name, by typing PHCM9795 in the "Directory name", and choose where you want to store the project by clicking the **Browse** button.



Step 4: Click **Create** to create your project.

You will now have a new folder in your directory, which contains only one file: PHCM9795.Rproj, and the two right-hand panes of RStudio will appear as below:



The top-right menu bar is showing that you are working within the PHCM9795 project, and the bottom-right window is showing the contents of that window: the single PHCM9795.Rproj file. We will add some more files to this project later.

1.4 A simple R analysis

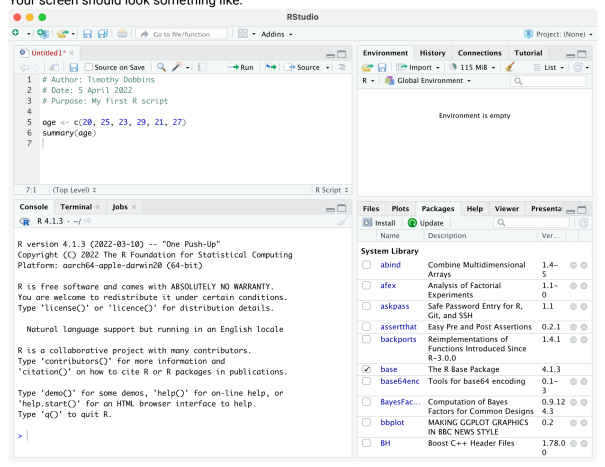
In this very brief section, we will introduce R by calculating the average of six ages.

To begin, open a new R Script by choosing **File > New file > R Script**. A script (or a program) is a collection of commands that are sequentially processed by R. You can also type Ctrl+Shift+N in Windows, or Command+Shift+N in MacOS to open a new script in RStudio, or click the **New File** button at the top of the RStudio window.

You should now see four window panes, as below. In the top-left window, type the following (replacing my name with yours, and including today's date):

```
# Author: Timothy Dobbins
# Date: 5 April 2022
# Purpose: My first R script
age <- c(20, 25, 23, 29, 21, 27)
summary(age)</pre>
```

Note: R is case-sensitive, so you should enter the text exactly as written in these notes. Your screen should look something like:



To run your script, choose **Code > Run Region > Run All**. You will see your code appear in the bottom-left window, with the following output:

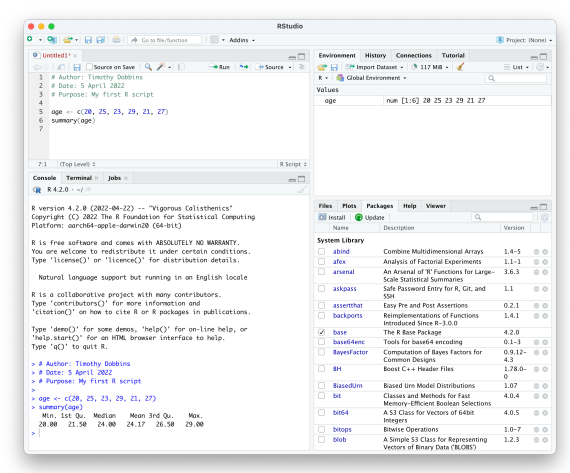
```
> # Author: Timothy Dobbins
> # Date: 5 April 2022
> # Purpose: My first R script
>
> age <- c(20, 25, 23, 29, 21, 27)
> summary(age)
    Min. 1st Qu. Median Mean 3rd Qu. Max.
    20.00 21.50 24.00 24.17 26.50 29.00
```

We will explain the key parts of this script later, but for now, you have entered six ages and calculated the mean age (along with five other summary statistics).

Save your script within the PHCM9795 project by using **File > Save As**, using the name my_first_analysis.R.

1.5 The RStudio environment

Now that we have seen a simple example of how to use R within RStudio, let's describe the RStudio environment. Let's assume that you have just run your first R script, and you have four windows as below:



The top-left window is call the **Source** window, and is where you write and edit your R scripts. Scripts can be saved by clicking **File > Save As** or by clicking on the symbol of a floppy disk at the top of the script. The file will have an extension of .R, for example script.R. Remember to give your script a meaningful title and remember to periodically save as you go.

In RStudio, the name of the script will be black when it has been saved, and will change to red if you have any unsaved changes.

The **Console** window, at the bottom left, contains the command line which is indicated with the symbol >. You can type commands here, but anything executed directly from the console is not saved and therefore is lost when the session ends (when you exit RStudio). You should always run your commands from a script file which you can save and use again later. When you run commands from a script, the output and any notes/errors are shown in the console. The Terminal and Jobs tabs will not be used in this course.

The **Environment** window at the top-right shows a list of objects that have been created during your session. When you close your RStudio session these objects will disappear. We will not use the History or Connections tabs in this course.

The bottom right corner contains some useful tabs, in particular the **Help** tab. When you are troubleshooting errors or learning how to use a function, the Help tab should be the first place you visit. Here you can search the help documents for all the packages you have installed. Whenever you create plots in R, these will be shown in the **Plots** tab. The **Packages** tab contains a list of installed

packages and indicates which ones are currently in use (we will learn about packages later). Packages which are loaded, i.e. in use, are indicated with a tick. Some packages are in use by default when you begin a new session. You can access information about a package by clicking on its name. The **Files** tab provides a shortcut to access your files. The Viewer tab will not be used in this course.

1.6 Some R basics

While we use R as a statistics package, R is a programming language. In order to use R effectively, we need to define some basics.

1.6.1 Scripts

While R can be run completely from the command line, issuing commands one-by-one, it is most commonly run using **scripts**. A script is simply a list of commands that are processed in order. The simple analysis we conducted earlier is a very simple script. Some things to know about R scripts:

- anything appearing after a # is a comment, and is ignored by R. The first three lines of our script are there for ourselves (either as writers of code, or readers of code). I include comments at the beginning of each of my scripts to describe:
 - who wrote the script (useful if someone else uses your script and wants to ask questions about it);
 - when the script was written;
 - what the script does. This last point may seem odd, but it's useful to describe what this script does, and why it might differ to other scripts being used in the analysis. This is particularly useful if your scripts become long and complex.
- R is case-sensitive. So age, AGE and Age could refer to three separate variables (please don't do this!)
- use blank lines and comments to separate sections of your script

1.6.2 Objects

If you do some reading about R, you may learn that R is an "object-oriented programming language". When we enter or import data into R, we are asking R to create **objects** from our data. These objects can be manipulated and transformed by **functions**, to obtain useful insights from our data. Objects in R are created using the **assignment operator**. The most common form of the assignment operator looks like an arrow: <- and is typed as the < and - symbols. The simplest way of reading <- is as the words "is defined as". Note that it possible to use -> and even = as assignment operators, but their use is less frequent.

Let's see an example:

```
x <- 42
```

This command creates a new object called x, which is defined as the number 42 (or in words, "x is defined as 42"). Running this command gives no output in the console, but the new object appears in the top-right **Environment** panel. We can view the object in the console by typing its name:

```
# Print the object x
x
```

```
## [1] 42
```

Now we see the contents of x in the console.

This example is rather trivial, and we rarely assign objects of just one value. In fact, we created an object earlier, called age, which comprised six values.

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1.6.3 Data structures

There are two main structures we will use to work with data in this course: **vectors** and **data frames**. A **vector** is a combination of data values, all of the same type. For example, our six ages that we entered earlier is a vector. You could think of a vector as a column of data (even though R prints vectors as rows!) And technically, even an object with only one value is a vector, a vector of size 1. The easiest way of creating a vector in R is by using the c() function, where c stands for 'combine'. In our previous Simple Analysis in R (Section 1.4), we wrote the command:

```
age <- c(20, 25, 23, 29, 21, 27)
```

This command created a new object called age, and *combined* the six values of age into one vector. Just as having a vector of size 1 is unusual, having just one column of data to analyse is also pretty unusual. The other structure we will describe here is a **data frame** which is essentially a collection of vectors, each of the same size. You could think of a data frame as being like a spreadsheet, with columns representing variables, and rows representing observations.

There are other structures in R, such as matrices and lists, which we won't discuss in this course. And you may come across the term **tibble**, which is a type of data frame.

1.6.4 Functions

If objects are the nouns of R, functions are the verbs. Essentially, functions transform objects. Functions can transform your data into summary statistics, graphical summaries or analysis results. For example, we used the summary() function to display summary statistics for our six ages. R functions are specified by their arguments (or inputs). The arguments that can be supplied for each function can be inspected by examining the help notes for that function. To obtain help for a function, we can submit help(summary) (or equivalently?summary) in the console, or we can use the **Help** tab in the bottom-right window of RStudio. For example, the first part of the help notes for summary appear as:

summary {base}

R Documentation

Object Summaries

Description

summary is a generic function used to produce result summaries of the results of various model fitting functions. The function invokes particular <u>methods</u> which depend on the <u>class</u> of the first argument.

Usage

The help notes in R can be quite cryptic, but the **Usage** section details what inputs should be specified for the function to run. Here, summary requires an object to be specified. In our case, we specified age, which is our object defined as the vector of six ages.

Most help pages also include some examples of how you might use the function. These can be found at the very bottom of the help page.

Examples

Run examples

```
summary(attenu, digits = 4) #-> summary.data.frame(...), default precision
summary(attenu $ station, maxsum = 20) #-> summary.factor(...)

lst <- unclass(attenu$station) > 20 # logical with NAs
## summary.default() for logicals -- different from *.factor:
summary(lst)
summary(as.factor(lst))
```

The summary() function is quite simple, in that it only requires one input, the object to be summarised. More complex functions might require a number of inputs. For example, the help notes for the descriptives() function in the jmv package show a large number of inputs can be specified:

| R Documentation | R Docume

Descriptives

Description

Descriptives are an assortment of summarising statistics, and visualizations which allow exploring the shape and distribution of data. It is good practice to explore your data with descriptives before proceeding to more formal tests.

Usage

```
descriptives(data, vars, splitBy = NULL, freq = FALSE,
  desc = "columns", hist = FALSE, dens = FALSE, bar = FALSE,
  barCounts = FALSE, box = FALSE, violin = FALSE, dot = FALSE,
  dotType = "jitter", boxMean = FALSE, boxLabelOutliers = TRUE,
  qq = FALSE, n = TRUE, missing = TRUE, mean = TRUE,
  median = TRUE, mode = FALSE, sum = FALSE, sd = TRUE,
  variance = FALSE, range = FALSE, min = TRUE, max = TRUE,
  se = FALSE, ci = FALSE, ciWidth = 95, iqr = FALSE,
  skew = FALSE, kurt = FALSE, sw = FALSE, pcEqGr = FALSE,
  pcNEqGr = 4, pc = FALSE, pcValues = "25,50,75", formula)
```

There are two things to note here. First, notice that the first two inputs are listed with no = symbol, but all other inputs are listed with = symbols (with values provided after the = symbol). This means that everything apart from data and vars have **default** values. We are free to not specify values for these inputs if we are happy with the defaults provided. For example, by default the variance is not calculated (as variance = FALSE). To obtain the variance as well as the standard deviation, we can change this default to variance = TRUE:

```
# Only the standard deviation is provided as the measure of variability
descriptives(data=pbc, vars=age)

# Additionally request the variance to be calculated
descriptives(data=pbc, vars=age, variance=TRUE)
```

Second, for functions with multiple inputs, we can specify the input name and its value, or we can ignore the input name and specify just the input values **in the order listed in the Usage section**. So the following are equivalent:

```
# We can specify that the dataset to be summarised is pbc,
# and the variable to summarise is age:
descriptives(data=pbc, vars=age)
# We can omit the input name, as long as we keep the inputs in the correct order -
```

1.6. SOME R BASICS

```
# that is, dataset first, variable second:
descriptives(pbc, age)

# We can change the order of the inputs, as long as we specify the input name:
descriptives(vars=age, data=pbc)
```

In this course, we will usually provide all the input names, even when they are not required. As you become more familiar with R, you will start to use the shortcut method.

1.6.4.1 The curse of inconsistency

As R is an open-source project, many people have contributed to its development. This has led to a frustrating part of R: some functions require a single object to be specified, but some require you to specify a data frame and select variables for analysis. Let's see an example.

The help for summary() specifies the usage as: summary(object, ...). This means we need to specify a single object to be summarised. An object could be a single column of data (i.e. a vector), or it could be a data frame. If we have a data frame called pbc which contains many variables, the command summary(pbc) would summarise every variable in the data frame.

What if we only wanted to summarise the age of the participants in the data frame? To select a single variable from a data frame, we can use the following syntax: dataframe\$variable. So to summarise just age from this data frame, we would use: summary(pbc\$age).

Compare this with the descriptives() function in the jmv package. We saw earlier that the two required inputs for descriptives() are data (the data frame to be analysed) and vars (the variables to be analysed). So to summarise age from the pbc data frame, we would specify descriptives(data=pbc, vars=age).

This inconsistency will seem maddening at first, and will continue to be maddening! Reading the **usage** section of the help pages is a useful way to determine whether you should specify an object (like pbc\$age) or a data frame and a list of variables.

1.6.5 Packages

A **package** is a collection of functions, documentation (and sometimes datasets) that extend the capabilities of R. Packages have been written by R users to be freely distributed and used by others. R packages can be obtained from many sources, but the most common source is CRAN: the Comprehensive R Archive Network.

A useful way of thinking about R is that R is like a smartphone, with packages being like apps which are downloaded from CRAN (similar to an app-store). When you first install R, it comes with a basic set of packages (apps) installed. You can do a lot of things with these basic packages, but sometimes you might want to do things differently, or you may want to perform some analyses that can't be done using the default packages. In these cases, you can install a package. Like installing an app on a smartphone, you only need to *install* a package once. But each time you want to use the package, you need to *load* the package into R.

1.6.6 How to install a package

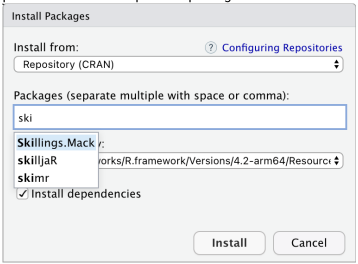
There are a couple of ways to install a package. You can use the install.packages() function if you know the exact name of the package. Let's use an example of installing the skimr package, which gives a very nice, high-level overview of any data frame. We can install skimr by typing the following into the console:

```
install.packages("skimr")
```

Note the use of the quotation marks.

Alternatively, RStudio offers a graphical way of installing packages that can be accessed via **Tools > Install Packages**, or via the **Install** button at the top of the **Packages** tab in the bottom-right window.

You can begin typing the name of the package in the dialog box that appears, and RStudio will use predictive text to offer possible packages:



While writing code is usually the recommended way to use R, installing packages is an exception. Using **Tools > Install Packages** is perfectly fine, because you only need to install a package once.

1.6.7 How to load a package

When you begin a new session in RStudio, i.e. when you open RStudio, only certain core packages are automatically loaded. You can use the library() function to load a package that you has previously been installed. For example, now that we have installed skimr, we need to load it before we can use it:

```
library(skimr)
```

Note that quotation marks are not required for the library() function (although they can be included if you really like quotation marks!).

Installing vs loading packages

Package installation:

- use the install.packages() function (note the 's') or Tools > Install packages
- the package name must be surrounded by quotation marks
- · only needs to be done once

Package loading

- use the library() function
- the package name does not need to be surrounded by quotation marks
- · must be done for each R session

1.7 What is this thing called the tidyverse?

If you have done much reading about R, you may have come across the tidyverse:

"The tidyverse is an opinionated collection of R packages designed for data science. All packages share an underlying design philosophy, grammar, and data structures." https://www.tidyverse.org/

Packages in the tidyverse have been designed with a goal to make using R more consistent by defining a "grammar" to manipulate data, examine data and draw conclusions from data. While the tidyverse is a common and powerful set of packages, we will not be teaching the tidyverse in this course for two main reasons:

- 1. The data we provide have been saved in a relatively tidy way, and do not need much manipulation for analyses to be conducted. The cognitive load in learning the tidyverse in this course is greater than the benefit that could be gained.
- 2. There are many resources (online, in print etc) that are based on base R, and do not use the tidyverse. It would be difficult to understand these resources if we taught only tidyverse techniques. In particular, the dataframe\$variable syntax is an important concept that should be understood before moving into the tidyverse.

In saying all of this, I think the tidyverse is an excellent set of packages, which I frequently use. At the completion of this course, you will be well equipped to teach yourself tidyverse using many excellent resources such as: Tidyverse Skills for Data Science and R for Data Science.

Part 2: Obtaining basic descriptive statistics

In this exercise, we will analyse data to complete a descriptive table from a research study. The data come from a study in primary biliary cirrhosis, a condition of the liver, from Modeling Survival Data: Extending the Cox Model Therneau and Grambsch [2010]. By the end of this exercise, we will have completed the following table.

Table 1.2: Summary of 418 partici	ipants from the PBC study ((Therneau and Grambsch.	. 2000)
-----------------------------------	-----------------------------	-------------------------	---------

Characteristic		Summary
Age (years)		Mean (SD) or Median [IQR]
Sex	Male	n (%)
Sex	Female	n (%)
AST* (U/ml)		Mean (SD) or Median [IQR]
Serum bilirubin		Mean (SD) or Median [IQR]
	1	n (%)
Stage	II	n (%)
	III	n (%)
	IV	n (%)
	Alive: no transplant	n (%)
Vital status at study end	Alive: transplant	n (%)
	Deceased	n (%)

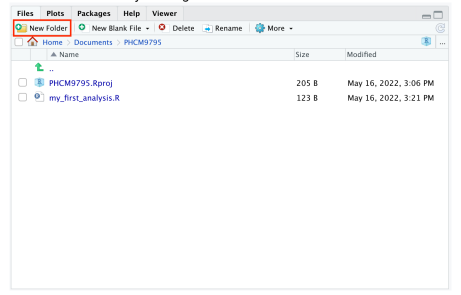
^{*} asparate aminotransferase

This table is available in Table1.docx, saved on Moodle.

1.8 Set up your data

We created a project in the previous step. We will now create a folder to store all the data for this course. Storing the data within the project makes life much easier!

Create a new folder by clicking the **New Folder** icon in the **Files** tab at the bottom-right:

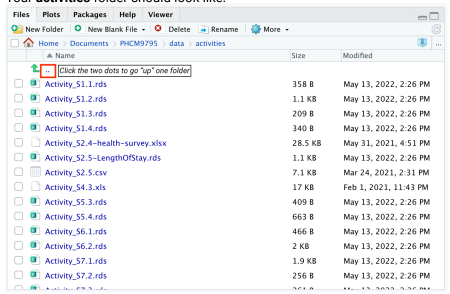


Call the new folder data.

Click on this folder to open it, and then create two new folders: activities and examples.

Download the "Data sets: for learning activities" from Moodle, and use Windows Explorer or MacOS Finder to save these data sets in **activities**. Save the "Data sets: example data from course notes" into the **examples** folder.

Your activities folder should look like:



Click the two dots next to the up-arrow at the top of the folder contents to move back up the folder structure. Note that you need to click the dots, and not the up-facing green arrow!

1.9 Reading a data file

Typing data directly into R is not common; we usually read data that have been previously saved. In this example, we will read an .rds file using the readRDS() function, which has only one input: the

location of the file.

- 1 Confirm that the pbc.rds file is in the activities sub-folder within the data folder (as per the previous steps).
- 2 Install the packages: jmv, skimr and summarytools using **Tools > Install packages**, or by typing into the console:

```
install.packages("jmv")
install.packages("skimr")
install.packages("summarytools")
```

3 - Load the skimr package, and use the readRDS() function to read the file into R, assigning it to a data frame called pbc. Because we set up our project, we can locate our data easily by telling R to use the file: "data/activities/pbc.rds", which translates as: the file pbc.rds which is located in the activities sub-folder within the data folder.

```
library(skimr)

pbc <- readRDS("data/activities/pbc.rds")</pre>
```

4 - We can now use the summary() function to examine the pbc dataset:

```
summary(pbc)
```

```
##
           'nг
                           time
                                          status
                                                              trt
##
    Min.
               1.0
                      Min.
                              : 41
                                      Min.
                                              :0.0000
                                                         Min.
                                                                :1.000
                      1st Qu.:1093
##
    1st Qu.:105.2
                                      1st Qu.:0.0000
                                                         1st Qu.:1.000
    Median :209.5
                     Median :1730
                                      Median :0.0000
                                                         Median :1.000
##
##
    Mean
            :209.5
                      Mean
                             :1918
                                      Mean
                                              :0.8301
                                                         Mean
                                                                :1.494
    3rd Ou.:313.8
                                      3rd Ou.:2.0000
##
                      3rd Ou.:2614
                                                         3rd Qu.:2.000
##
    Max.
            :418.0
                             :4795
                                              :2.0000
                                                                 :2.000
                      Max.
                                      Max.
                                                         Max.
##
                                                         NA's
                                                                :106
##
                                          ascites
                                                               hepato
          age
                           sex
##
    Min.
            :26.28
                      Min.
                             :1.000
                                       Min.
                                               :0.00000
                                                           Min.
                                                                   :0.0000
    1st Qu.:42.83
                                       1st Qu.:0.00000
##
                      1st Qu.:2.000
                                                           1st Qu.:0.0000
##
    Median :51.00
                     Median :2.000
                                       Median :0.00000
                                                           Median :1.0000
##
    Mean
            :50.74
                      Mean
                             :1.895
                                       Mean
                                               :0.07692
                                                           Mean
                                                                   :0.5128
    3rd Qu.:58.24
                      3rd Qu.:2.000
                                       3rd Qu.:0.00000
                                                           3rd Qu.:1.0000
##
##
    Max.
            :78.44
                      Max.
                              :2.000
                                       Max.
                                               :1.00000
                                                           Max.
                                                                   :1.0000
                                                                   :106
##
                                       NA's
                                               :106
                                                           NA's
##
       spiders
                           edema
                                               bili
                                                                 chol
                                                 : 0.300
                              :0.0000
##
    Min.
            :0.0000
                       Min.
                                         Min.
                                                            Min.
                                                                    : 120.0
##
    1st Qu.:0.0000
                       1st Qu.:0.0000
                                         1st Qu.: 0.800
                                                            1st Qu.: 249.5
                       Median :0.0000
##
    Median :0.0000
                                         Median : 1.400
                                                            Median : 309.5
    Mean
            :0.2885
                       Mean
                               :0.1005
                                         Mean
                                                 : 3.221
                                                            Mean
                                                                    : 369.5
##
    3rd Ou.:1.0000
                       3rd Qu.:0.0000
                                         3rd Qu.: 3.400
                                                            3rd Ou.: 400.0
                                                 :28.000
##
    Max.
            :1.0000
                       Max.
                               :1.0000
                                         Max.
                                                            Max.
                                                                    :1775.0
            :106
##
    NA's
                                                            NA's
                                                                    :134
##
       albumin
                          copper
                                            alkphos
                                                                 ast
##
    Min.
            :1.960
                      Min.
                             : 4.00
                                        Min.
                                                :
                                                   289.0
                                                            Min.
                                                                    : 26.35
##
    1st Qu.:3.243
                      1st Qu.: 41.25
                                        1st Qu.: 871.5
                                                            1st Qu.: 80.60
##
    Median :3.530
                      Median : 73.00
                                        Median : 1259.0
                                                            Median :114.70
##
    Mean
            :3.497
                             : 97.65
                                                : 1982.7
                                                                    :122.56
                      Mean
                                        Mean
                                                            Mean
##
    3rd Qu.:3.770
                      3rd Qu.:123.00
                                        3rd Qu.: 1980.0
                                                            3rd Qu.:151.90
##
                                                :13862.4
    Max.
            :4.640
                      Max.
                             :588.00
                                        Max.
                                                            Max.
                                                                    :457.25
##
                      NA's
                             :108
                                        NA's
                                                :106
                                                            NA's
                                                                    :106
##
                          platelet
                                           protime
          trig
                                                              stage
```

```
##
           : 33.00
                            : 62.0
                                            : 9.00
   Min.
                     Min.
                                     Min.
                                                     Min.
                                                             :1.000
##
   1st Qu.: 84.25
                     1st Qu.:188.5
                                     1st Qu.:10.00
                                                     1st Qu.:2.000
  Median :108.00
                     Median :251.0
                                                     Median :3.000
##
                                     Median :10.60
## Mean
           :124.70
                     Mean
                            :257.0
                                     Mean
                                            :10.73
                                                     Mean
                                                             :3.024
  3rd Qu.:151.00
                     3rd Qu.:318.0
                                     3rd Qu.:11.10
                                                     3rd Qu.:4.000
##
           :598.00
                            :721.0
                                     Max.
                                            :18.00
                                                             :4.000
## Max.
                     Max.
                                                     Max.
##
  NA's
           :136
                     NA's
                            :11
                                     NA's
                                            :2
                                                     NA's
                                                             :6
```

An alternative to the summary() function is the skim() function in the skimr package, which produces summary statistics as well as rudimentary histograms:

skim(p	pbc)									
— Data	Summary —									
	•	Values								
Name		pbc								
Number	of rows	418								
Number	of columns	20								
Column	type frequency:									
numer		20								
Group v	ariables	None								
Vosi	oblo tupo, pumorio									
	able type: numeric _variable n_missing		mean	sd	p0	p25	p50	p75	n100	hist
1 id	_vaitabte ii_iii1331ii	–	210.	121.	1	105.	210.	314.	418	11131
2 time	(1918.	1105.	41	1093.	<u>1</u> 730	2614.	4795	=
3 stati			0.830	0.956	0	<u>1</u> 033.	0	2	<u>4</u> 733	
4 trt	106		1.49	0.501	1	1	1	2	2	П
5 age	100		50.7	10.4	26.3	42.8	51.0	58.2	78.4	
6 sex			1.89	0.307	1	2	2	2	2	
7 asci			0.0769		0	0	0	0	1	
8 hepa			0.513	0.501	0	0	1	1	1	
9 spide			0.288	0.454	0	0	0	1	1	
10 edema) 1	0.100	0.253	0	0	0	0	1	
11 bili	() 1	3.22	4.41	0.3	0.8	1.4	3.4	28	
12 chol	134	0.679	370.	232.	120	250.	310.	400	<u>1</u> 775	
13 albu	min () 1	3.50	0.425	1.96	3.24	3.53	3.77	4.64	
14 coppe	er 108	0.742	97.6	85.6	4	41.2	73	123	588	
15 alkpl		0.746	<u>1</u> 983.	<u>2</u> 140.	289	872.	<u>1</u> 259	<u>1</u> 980	<u>13</u> 862.	
16 ast	106	0.746	123.	56.7	26.4	80.6	115.	152.	457.	
17 trig	136	0.675	125.	65.1	33	84.2	108	151	598	
18 plate	elet 13	0.974	257.	98.3	62	188.	251	318	721	
19 prot	ime 2	0.995	10.7	1.02	9	10	10.6	11.1	18	
20 stage	е (0.986	3.02	0.882	1	2	3	4	4	

The summary() and skim() functions are useful to give a quick overview of a dataset: how many variables are included, how variables are coded, which variables contain missing data and a crude histogram showing the distribution of numeric variables.

1.10 Summarising continuous variables

One of the most flexible functions for summarising continuous variables is the descriptives() function from the jmv package. The function is specified as descriptives(data=, vars=) where:

- data specifies the dataframe to be analysed
- vars specifies the variable(s) of interest, with multiple variables combined using the c() function

We can summarise the three continuous variables in the pbc data: age, AST and serum bilirubin, as shown below.

```
library(jmv)
descriptives(data=pbc, vars=c(age, ast, bili))
##
##
    DESCRIPTIVES
##
##
    Descriptives
##
##
                                                       bili
                              age
                                           ast
##
##
      Ν
                                   418
                                                312
                                                              418
##
      Missing
                                     0
                                                106
                                                                a
##
      Mean
                              50.74155
                                           122.5563
                                                         3.220813
                              51.00068
##
      Median
                                           114.7000
                                                         1.400000
                              10.44721
##
      Standard deviation
                                           56.69952
                                                         4.407506
##
      Minimum
                              26.27789
                                           26.35000
                                                        0.3000000
##
      Maximum
                              78.43943
                                           457.2500
                                                         28.00000
##
```

By default, the descriptives function presents the mean, median, standard deviation, minimum and maximum. We can request additional statistics, such as the quartiles (which are called the percentiles, or pc, in the descriptives function):

```
descriptives(data=pbc, vars=c(age, ast, bili), pc=TRUE)
```

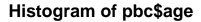
```
##
    DESCRIPTIVES
##
##
    Descriptives
##
##
                                                       bili
                              age
                                           ast
##
##
                                   418
                                                              418
      Ν
                                                312
##
      Missing
                                     0
                                                106
                                                                0
##
      Mean
                              50.74155
                                           122.5563
                                                         3.220813
##
      Median
                              51.00068
                                           114.7000
                                                        1.400000
##
      Standard deviation
                              10.44721
                                           56.69952
                                                         4.407506
##
      Minimum
                              26.27789
                                           26.35000
                                                        0.3000000
##
      Maximum
                              78.43943
                                           457.2500
                                                        28.00000
##
      25th percentile
                              42.83231
                                           80.60000
                                                        0.8000000
##
      50th percentile
                              51.00068
                                           114.7000
                                                        1.400000
##
      75th percentile
                              58.24093
                                           151.9000
                                                        3.400000
##
```

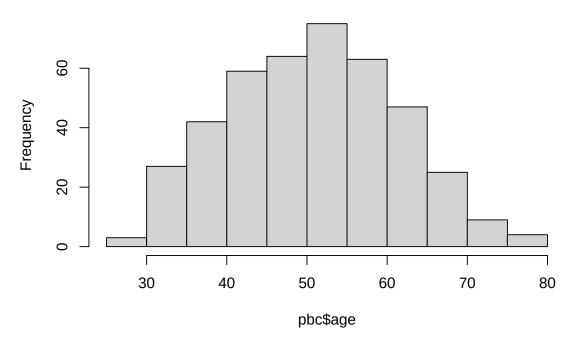
1.11 Producing a histogram

##

We can use the hist() function to produce a histogram, specifying the dataframe to use and the variable to be plotted as dataframe\$variable:

```
hist(pbc$age)
```

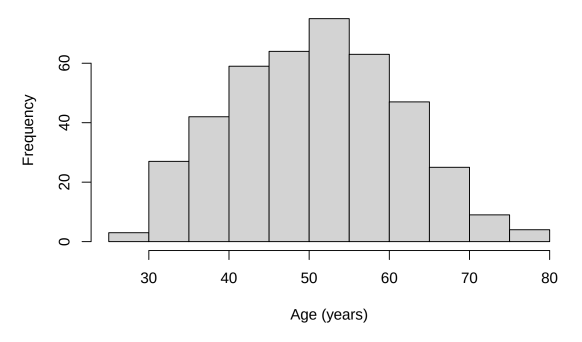




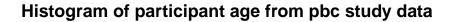
The histogram function does a remarkably good job of choosing cutpoints and binwidths, and these rarely need to be changed. However, the labelling of the histogram should be improved by using xlab="" and main="" to assign labels for the x-axis and overall title respectively:

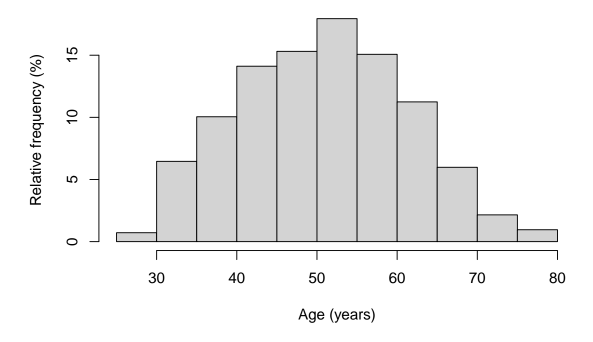
```
hist(pbc$age, xlab="Age (years)",
    main="Histogram of participant age from pbc study data")
```

Histogram of participant age from pbc study data



By default, the hist() function plots a **frequency histogram**, with counts on the y-axis. We can tweak the histogram using the following code to plot a histogram of the **relative frequencies**:



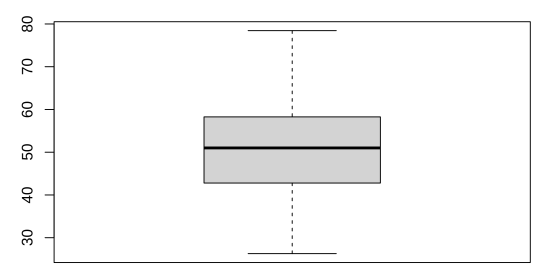


1.12 Producing a boxplot

The boxplot function is used to produce boxplots, again specifying the dataframe to use and the variable to be plotted as dataframe\$variable. Labels can be applied in the same way as the histogram:

boxplot(pbc\$age, xlab="Age (years)", main="Boxplot of participant age from pbc study data")





Age (years)

1.13 Producing a one-way frequency table

We have three categorical variables to summarise in Table 1: sex, stage and vital status. These variables are best summarised using one-way frequency tables.

```
library(summarytools)
freq(pbc$sex)
## Frequencies
## pbc$sex
## Type: Numeric
##
##
                      % Valid
                               % Valid Cum.
                                            % Total
                                                     % Total Cum.
               Freq
##
                       _____ ____
##
                 44
           1
                       10.53
                                     10.53
                                              10.53
                                                            10.53
##
                                              89.47
           2
                374
                       89.47
                                   100.00
                                                           100.00
##
         <NA>
                  0
                                               0.00
                                                           100.00
##
        Total
                418
                      100.00
                                    100.00
                                             100.00
                                                           100.00
```

1.13.1 Defining categorical variables as factors

You will notice that the table above, in its current form, is uninterpretable as the 1 and 2 categories are not labelled. In this course, all variables including categorical variables tend to be numerically coded. To define a categorical variable as such in R, we define it as a **factor** using the factor function: factor(variable=, levels=, labels=)
We specify:

- · levels: the values the categorical variable can take
- · labels: the labels corresponding to each of the levels (entered in the same order as the levels)

To define our variable sex as a factor, we use:

```
pbc$sex <- factor(pbc$sex, levels=c(1, 2), labels=c("Male", "Female"))</pre>
```

We can confirm the coding by re-running a frequency table:

```
freq(pbc$sex)
## Frequencies
## pbc$sex
## Type: Factor
##
##
                   % Valid % Valid Cum. % Total % Total Cum.
## ----- --- ----
##
        Male
              44
                    10.53
                                10.53
                                       10.53
                                                   10.53
              374
##
                   89.47
                               100.00
                                       89.47
                                                  100.00
      Female
##
        <NA>
              0
                                        0.00
                                                  100.00
                 100.00
##
       Total
              418
                               100.00
                                       100.00
                                                  100.00
```

Task: define stage and status (Vital Status) as factors, and produce one-way frequency tables. Refer to the file pbc_info.txt to view the labels for each variable. For example, for Stage:

```
pbcstage <- factor(pbcstage, levels=c(1, 2, 3, 4), labels=c("Stage 1", "Stage 2", "Stage 3", "Stage freq(pbcstage)
```

```
## pbc$stage
## Type: Factor
##
            Freq % Valid % Valid Cum. % Total % Total Cum.
##
##
 ##
     Stage 1
            21
                  5.10
                            5.10
                                   5.02
                                             5.02
                  22.33
                           27.43
##
     Stage 2
             92
                                  22.01
                                            27.03
##
     Stage 3
           155 37.62
                           65.05 37.08
                                            64.11
                 34.95
     Stage 4 144
                          100.00 34.45
##
                                            98.56
##
       <NA>
             6
                                  1.44
                                           100.00
                                  100.00
                      100.00
##
       Total
             418 100.00
                                            100.00
```

1.14 Producing a two-way frequency table

Frequencies

To produce tables summarising two categorical variables, we can use the contTables() function within the jmv package. The minimal inputs to include are data: the name of the data frame to be analysed, rows: the variable representing the rows of the table, and cols: the name of the columns of the table.

For example, to produce a two-way table showing stage of disease by sex using the pbc data frame, we use:

```
contTables(data=pbc, rows=sex, cols=stage)
```

```
##
##
    CONTINGENCY TABLES
##
##
    Contingency Tables
##
##
                   Stage 1
                                Stage 2
                                             Stage 3
                                                          Stage 4
                                                                       Total
       sex
##
       Male
                          3
                                       8
                                                                17
                                                                           44
##
                                                   16
                                                               127
##
       Female
                         18
                                      84
                                                  139
                                                                          368
       Total
                         21
                                      92
                                                  155
                                                               144
                                                                          412
##
##
##
##
##
    x<sup>2</sup> Tests
##
##
              Value
                              df
                                     p
##
##
       χ²
              0.8779873
                               3
                                     0.8307365
##
       N
                     412
##
```

[The bottom part of the output, χ^2 Tests, can be ignored for now]

You may notice in the above that the number of observations is now 412. This is because there are missing observations for either sex or stage: which is it, and how would you determine this? From the cross-tabulation, you can see the individual frequencies of participants in each of the categories in each cell. For example, there are 3 male participants who have Stage 1 disease. You can also read the totals for each row and column. For example, there are 44 males, and 144 participants have Stage 4 disease.

You can also add percentages into your table using pcCol=TRUE to include column percents, and pcRow=TRUE for row percents. For example, to calculate the relative frequencies (i.e. percentages) of sex within each stage, we would request **column percents** with the option: pcCol=TRUE.

```
contTables(data=pbc, rows=sex, cols=stage, pcCol=TRUE)
##
##
    CONTINGENCY TABLES
##
##
    Contingency Tables
##
##
                                 Stage 1
                                              Stage 2
                                                                         Stage 4
                                                                                      Total
      sex
                                                            Stage 3
##
##
               Observed
     Male
                                                                 16
                                                                             17
##
            % within column
                                14.28571
                                              8.69565
                                                         10.32258
                                                                      11.80556
                                                                                   10.67961
##
##
     Female
                Observed
                                         18
                                                     84
                                                                139
                                                                            127
                                                                                        368
            % within column
                                85.71429
                                             91.30435
##
                                                          89.67742
                                                                      88.19444
                                                                                   89.32039
##
##
     Total
                Observed
                                        21
                                                     92
                                                                155
                                                                            144
                                                                                        412
##
            % within column
                               100.00000
                                            100.00000
                                                         100.00000
                                                                       100.00000
                                                                                    100.00000
##
##
##
##
    x<sup>2</sup> Tests
##
##
             Value
                            df
                                  р
```

```
##
## x<sup>2</sup> 0.8779873 3 0.8307365
## N 412
##
```

We can see that the 3 male participants with Stage 1 disease represent 14% of those with Stage 1 disease.

1.15 Saving data in R

There are many ways to save data from R, depending on the type of file you want to save. The recommendation for this course is to save your data using the .rds format, using the saveRDS() function, which takes two inputs: saveRDS(object, file). Here, object is the R object to be saved (usually a data frame), and file is the location for the file to be saved (file name and path, including the .rds suffix).

It is not necessary to save our PBC data, as we have made only minor changes to the data that can be replicated by rerunning our script. If you had made major changes and wanted to save your data, you could use:

```
saveRDS(pbc, file="pbc_revised.rds")
```

1.16 Copying output from R

It is important to note that saving your data or your script in R will not save your output. The easiest way to retain the output of your analyses is to copy the output from the Console into a word processor package (e.g. Microsoft Word) before closing R.

Unfortunately, by default, R is not ideal for creating publication quality tables. There are many packages that will help in this process, such as R Markdown, bookdown¹, huxtable, gt and gtsummary, but their use is beyond the scope of this course. R Markdown for Scientists provides an excellent introduction to R Markdown.

Task: Complete Table 1 using the output generated in this exercise. You should decide on whether to present continuous variables by their means or medians, and present the most appropriate measure of spread. Include footnotes to indicate if any variables contain missing observations.

Part 3: Creating other types of graphs

The plot() function, also known as base graphics, is the default method of plotting data in R that can produce publication-quality graphics with minimal coding. There are alternative packages for plotting, with ggplot2 being one of the most well known. We will present instructions for base graphics in this course, but excellent documentation for ggplot2 can be found at the ggplot2: Elegant Graphics for Data Analysis website, written by the package authors.

1.17 Bar graphs

The simplest way to use the plot() function is by specifying an object to be plotted. As with the hist() function, to plot a single variable from a data frame, we must define it using: dataframe\$variable.

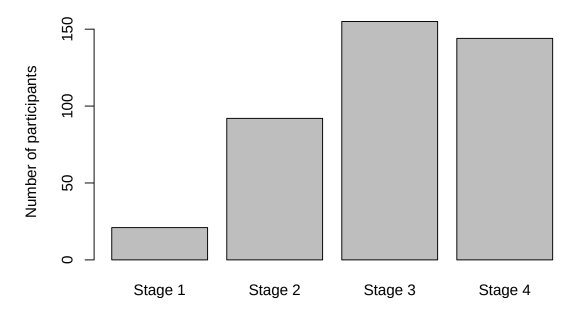
Here we will create the bar chart shown in Figure 1.3 of the statistics notes using the pbc.rds dataset. The x-axis of this graph will be the stage of disease, and the y-axis will show the number of participants in each category.

¹these R notes and the PHCM9795 course notes have been written using bookdown

1.17. BAR GRAPHS 31

```
plot(pbc$stage,
    main="Bar graph of stage of disease from PBC study",
    ylab="Number of participants")
```

Bar graph of stage of disease from PBC study



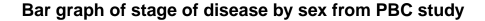
Note that stage is a categorical variable, that has been defined as a factor (in Section 1.13.1). You **must define categorical data as factors** to plot them in a bar graph.

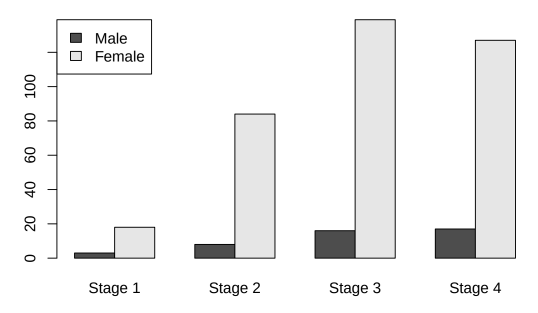
1.17.1 Clustered bar graph

To create a clustered bar chart as shown in Figure 1.4 of the statistics notes, we need to do a bit of manipulation. We first need to tabulate the data using the table() function. We want to plot stage of disease broken down by sex, so we specify sex as the first variable, and stage as the second variable for the table() command.

```
counts <- table(pbc$sex, pbc$stage)</pre>
counts
##
##
             Stage 1 Stage 2 Stage 3 Stage 4
##
     Male
                    3
                             8
                                    16
                                             17
     Female
                   18
                            84
                                   139
                                             127
```

After tabulating the data, we use the barplot() function to plot the summarised data. We specify the main title using main=" ", specify that the stages be plotted separately by sex (beside=TRUE), specify the legend be defined by sex, and position the legend in the top-left of the graph:



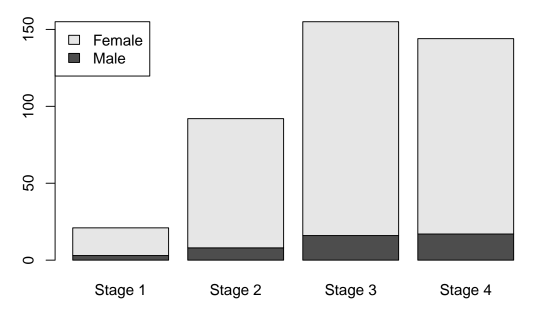


1.17.2 Stacked bar graph

A stacked bar graph can be constructed as for the clustered bar graph, but we specify beside=FALSE:

1.17. BAR GRAPHS 33





1.17.3 Stacked bar graph of relative frequencies

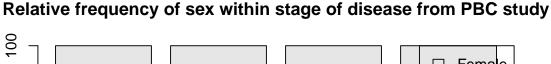
To plot relative frequencies, we need to transform our table of frequencies (counts) into proportions, by using the prop.table() function. The prop.table() function takes two arguments: a table of counts, and margin, which defines whether we want proportions calculated by row (margin=1) or column (margin=2).

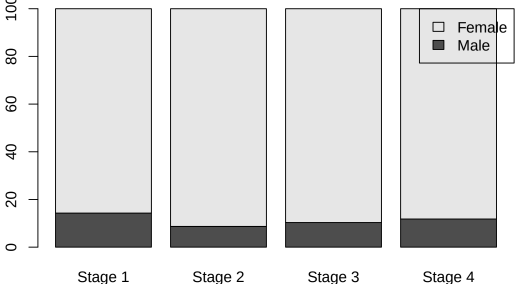
We want to calculate the relative frequency of sex within each stage category. From our counts table above, this equates to calculating *column* proportions, so we specify margin=2. We also multiply the resulting table by 100 to obtain percentages (rather than proportions):

```
percent <- prop.table(counts, margin=2)*100
percent

##
## Stage 1 Stage 2 Stage 3 Stage 4
## Male 14.285714 8.695652 10.322581 11.805556
## Female 85.714286 91.304348 89.677419 88.194444</pre>
```

After calculating the percentages, we use barplot() again, similar to the stacked bar graph:





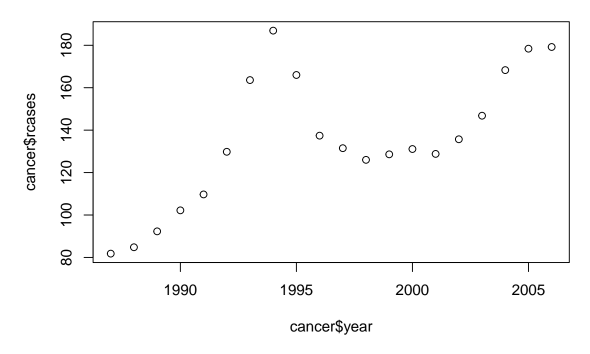
1.18 Creating line graphs

To demonstrate the graphing of aggregate data, we use the data on new cases and deaths from prostate cancer in males in NSW. This data has been entered as Example_1.2.rds.

```
cancer <- readRDS("data/examples/Example_1.2.rds")</pre>
summary(cancer)
                                       ndeaths
                                                                          rdeaths
##
         year
                        ncases
                                                          rcases
##
    Min.
           :1987
                    Min.
                           :1567
                                    Min.
                                           : 645.0
                                                      Min.
                                                             : 81.8
                                                                       Min.
                                                                              :31.10
    1st Qu.:1992
                                    1st Qu.: 788.2
                                                      1st Qu.:121.9
                                                                       1st Qu.:34.67
##
                    1st Qu.:2804
    Median :1996
                   Median :3790
                                   Median : 868.0
                                                      Median :131.3
                                                                       Median :36.55
##
##
    Mean
           :1996
                    Mean
                           :3719
                                    Mean
                                           : 855.0
                                                             :135.4
                                                                       Mean
                                                                              :37.09
                                                      Mean
##
    3rd Qu.:2001
                    3rd Qu.:4403
                                    3rd Qu.: 921.0
                                                      3rd Qu.:164.2
                                                                       3rd Qu.:40.38
##
    Max.
           :2006
                   Max.
                           :6158
                                    Max.
                                           :1044.0
                                                      Max.
                                                             :186.9
                                                                       Max.
                                                                              :43.80
```

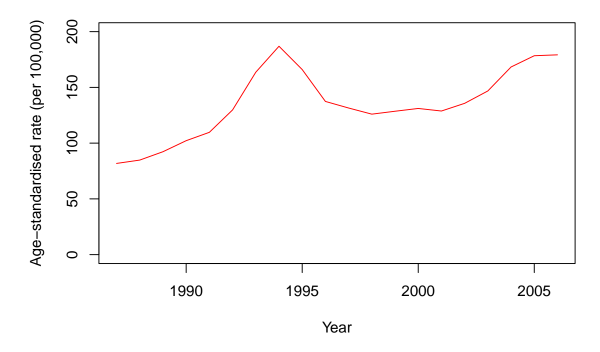
We begin by plotting cancer cases (as the *y* variable) against year (the *x* variable).

```
plot(x=cancer$year, y=cancer$rcases)
```

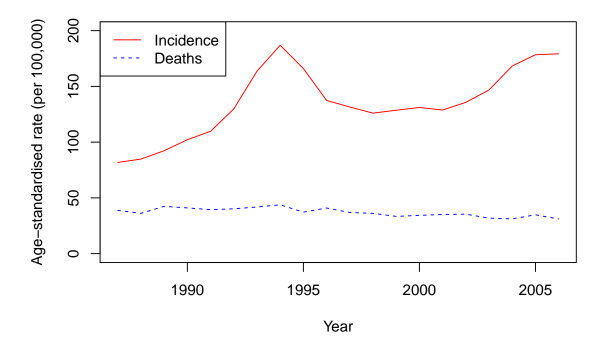


Let's define the plot to be joined by lines (type="1"), in the colour red (col="red"), providing meaningful labels for the x-axis and y-axis, and changing the scale of the y-axis to be between 0 and 200 (ylim=c(0,200)):

```
plot(x=cancer$year, y=cancer$rcases,
    type="l", col = "red",
    xlab = "Year",
    ylab = "Age-standardised rate (per 100,000)", ylim=c(0,200))
```



We can now add a second line to the plot using the lines() function, specifying a dashed line (lty=2), and add a legend to the plot:



Note: coding for graphs is not always straightforward. Two excellent resources for creating graphs in R are: R Graphics Cookbook and The R Graph Gallery.

Module 2

Probability and probability distributions

2.1 Importing data into R

We have described previously how to import data that have been saved as R .rds files. It is quite common to have data saved in other file types, such as Microsoft Excel, or plain text files. In this section, we will demonstrate how to import data from other packages into R.

There are two useful packages for importing data into R: haven (for data that have been saved by Stata, SAS or SPSS) and readx1 (for data saved by Microsoft Excel). Additionally, the labelled package is useful in working with data that have been labelled in Stata.

2.1.1 Importing plain text data into R

A csv file, or a "comma separated variables" file is commonly used to store data. These files have a very simple structure: they are plain text files, where data are separated by commas. csv files have the advantage that, as they are plain text files, they can be opened by a large number of programs (such as Notepad in Windows, TextEdit in MacOS, Microsoft Excel - even Microsoft Word). While they can be opened by Microsoft Excel, they can be opened by many other programs: the csv file can be thought of as the lingua-franca of data.

In this demonstration, we will use data on the weight of 1000 people entered in a csv file called weight_s2.csv available on Moodle.

To confirm that the file is readable by any text editor, here are the first ten lines of the file, opened in Notepad on Microsoft Windows, and TextEdit on MacOS.



We can use the read.csv function:

```
sample <- read.csv("data/examples/Weight_s2.csv")</pre>
```

Here, the read.csv function has the default that the first row of the dataset contains the variable names. If your data do not have column names, you can use header=FALSE in the function.

Note: there is an alternative function read_csv which is part of the readr package (a component of the tidyverse). Some would argue that the read_csv function is more appropriate to use because of an issue known as strings.as.factors. The strings.as.factors default was removed in R Version 4.0.0, so it is less important which of the two functions you use to import a .csv file. More information about this issue can be found here and here.

2.2 Checking your data for errors in R

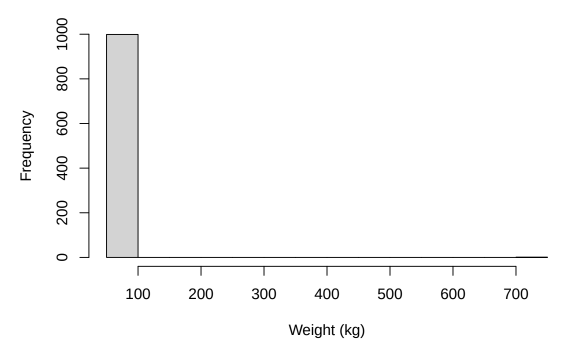
Before you start describing and analysing your data, it is important to make sure that no errors have been made during the data entry process. Basically, you are looking for values that are outside the range of possible or plausible values for that variable.

If an error is found, the best method for correcting the error is to go back to the original data e.g. the hard copy questionnaire, to obtain the original value, entering the correct value into R If the original data is not available or the original data is also incorrect, the erroneous value is often excluded from the dataset.

For continuous variables, the easiest methods are to examine a boxplot and histogram. For example, a boxplot and histogram for the weight variable we just imported appear as:

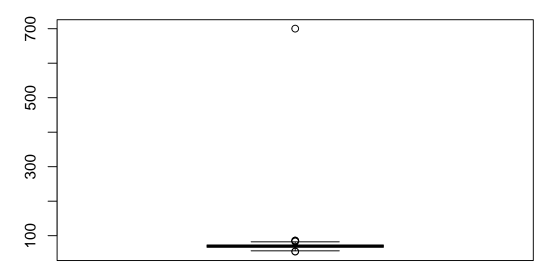
hist(sample\$weight, xlab="Weight (kg)", main="Histogram of 1000 weights")

Histogram of 1000 weights



boxplot(sample\$weight, xlab="Weight (kg)", main="Boxplot of 1000 weights")

Boxplot of 1000 weights



Weight (kg)

There is a clear outlying point shown in the boxplot. Although not obvious, the same point is shown in the histogram as a bar around 700 with a very short height.

We can identify any outlying observations in the dataset using the subset function. You will need to decide if these values are a data entry error or are biologically plausible. If an extreme value or "outlier", is biologically plausible, it should be included in all analyses.

For example, to list any observations from the sample dataset with a weight larger than 200:

```
subset(sample, weight>200)
## id weight
```

We see that there is a very high value of 700.2kg. A value as high as 700kg is likely to be a data entry error (e.g. error in entering an extra zero) and is not a plausible weight value. Here, **you should check your original data**.

You might find that the original weight was recorded in medical records as 70.2kg. You can change this in R by writing code.

Note: many statistical packages will allow you to view a spreadsheet version of your data and edit values in that spreadsheet. This is not best practice, as corrected observations may revert to their original values depending on whether the edited data have been saved or not. By using code-based recoding, the changes will be reproduced the next time the code is run.

We will use an ifelse statement to recode the incorrect weight of 700.2kg into 70.2kg. The form of the ifelse statement is as follows:

```
ifelse(test, value_if_true, value_if_false)
```

Our code will create a new column (called weight_clean) in the sample dataframe. We will test whether weight is equal to 700.2; if this is true, we will assign weight_clean to be 70.2, otherwise weight_clean will equal the value of weight.

Putting it all together:

58 58 700.2

```
sample$weight_clean = ifelse(sample$weight==700.2, 70.2, sample$weight)
```

Note: if an extreme value lies within the range of biological plausibility it should not be removed from analysis.

Once you have checked your data for errors, you are ready to start analysing your data.

2.2.1 What on earth: == ?

In R, the test of equality is denoted by two equal signs: ==. So we would use == to test whether an observation is equal to a certain value. Let's see an example:

```
# Test whether 6 is equal to 6
6 == 6
## [1] TRUE

# Test whether 6 is equal to 42
6 == 42
## [1] FALSE
```

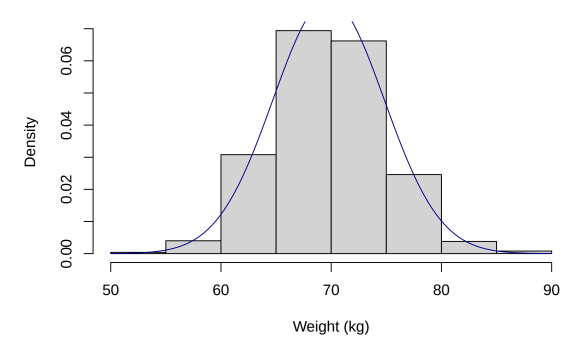
You can read the == as "is equal to". So the code sample\$weight == 700.2 is read as: "is the value of weight from the data frame sample equal to 700.2?". In our ifelse statement above, if this condition is true, we replace weight by 70.2; if it is false, we leave weight as is.

2.3 Overlaying a Normal curve on a histogram

It can be useful to produce a histogram with an overlayed Normal curve to assess whether our sample appears approximately Normally distributed. We can do this by plotting a histogram using the hist() function. As we're overlaying a probability distribution, we request the histogram be plotted on a probability scale, rather than a frequency scale, using probability=TRUE. We then request a curve be overlayed using the curve() function:

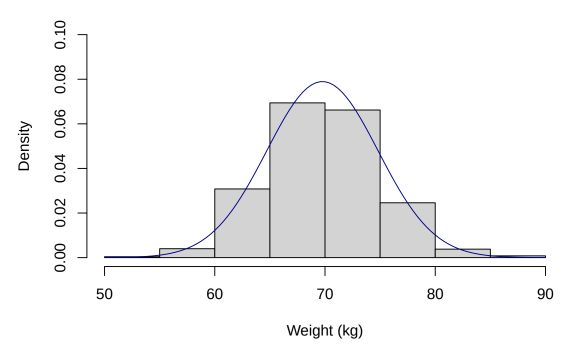
- the curve should be based on the Normal distribution (dnorm);
 - with a mean equal to the mean of the cleaned weight: mean(sample\$weight_clean));
 - and a standard deviation equal to the standard deviation of the cleaned weight: sd(sample\$weight_clean))
- · using a dark-blue colour;
- and added to the previous histogram (rather than plotting the curve by itself): add=TRUE

Histogram of 1000 weights



Notice that the top of the curve is chopped off. We can plot the whole curve by extending the y-axis of the histogram to 0.1:





2.4 Descriptive statistics for checking normality

All the descriptive statistics including *skewness* and *kurtosis* discussed in this module can be obtained using the descriptives function from the jmv package. In particular, skewness and kurtosis can be requested in addition to the default statistics by including: skew=TRUE, kurt=TRUE:

```
library(jmv)

descriptives(data=sample, vars=weight_clean, skew=TRUE, kurt=TRUE)
```

```
##
##
    DESCRIPTIVES
##
    Descriptives
##
##
##
                               weight_clean
##
##
                                        1000
##
      Missing
##
      Mean
                                    69.76450
      Median
                                    69.80000
##
##
      Standard deviation
                                    5.052676
##
      Minimum
                                    53.80000
##
      Maximum
                                    85.80000
##
      Skewness
                                 0.07360659
##
      Std. error skewness
                                 0.07734382
##
      Kurtosis
                                 0.05418774
##
      Std. error kurtosis
                                  0.1545343
```

##

2.5 Importing Excel data into R

Another common type of file that data are stored in is a Microsoft Excel file (.xls or .xlsx). In this demonstration, we will import a selection of records from a large health survey, stored in the file health-survey.xlsx.

The health survey data contains 1140 records, comprising:

- sex: 1 = respondent identifies as male; 2 = respondent identifies as female
- · height: height in meters
- weight: weight in kilograms

To import data from Microsoft Excel, we can use the read_excel() function in the readxl package.

```
library(readxl)
survey <- read_excel("data/examples/health-survey.xlsx")
summary(survey)</pre>
```

```
##
                      height
                                     weight
        sex
##
  Min. :1.00
                  Min. :1.220
                                 Min. : 22.70
                                 1st Qu.: 68.00
##
   1st Qu.:1.00
                  1st Qu.:1.630
##
   Median :2.00
                  Median :1.700
                                 Median : 79.40
## Mean
         :1.55
                  Mean :1.698
                                 Mean
                                        : 81.19
## 3rd Qu.:2.00
                  3rd Qu.:1.780
                                 3rd Qu.: 90.70
## Max.
         :2.00
                         :2.010
                                        :213.20
                  Max.
                                 Max.
```

We can see that sex has been entered as a numeric variable. We should transform it into a factor so that we can assign labels to each category:

```
survey$sex <- factor(survey$sex, level=c(1,2), labels=c("Male", "Female"))
summary(survey$sex)</pre>
```

```
## Male Female
## 513 627
```

We also note that height looks like it has been entered as meters, and weight as kilograms.

2.6 Generating new variables

Our health survey data contains information on height and weight. We often summarise body size using BMI: body mass index which is calculated as: $\frac{\text{weight (kg)}}{(\text{height (m)})^2}$

We can create a new column in our data frame in many ways, such as using the following approach: dataframe\$new_column <- <formula> For example:

```
survey$bmi <- survey$weight / (survey$height^2)</pre>
```

We should check the construction of the new variable by examining some records. The head() and tail() functions list the first and last 6 records in any dataset. We can also examine a histogram and boxplot:

head(survey)

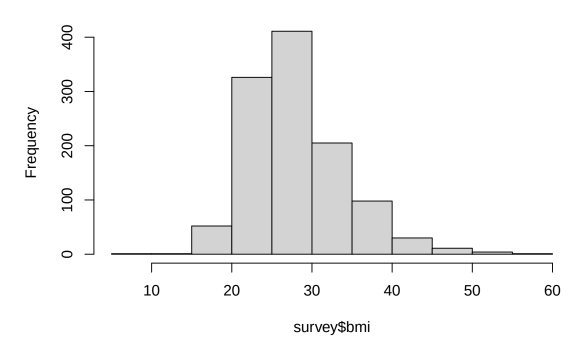
```
## # A tibble: 6 x 4
            height weight
##
     sex
            <dbl>
                    <dbl> <dbl>
##
     <fct>
             1.63
                     81.7 30.8
## 1 Male
## 2 Male
              1.63
                     68
                           25.6
                     97.1
## 3 Male
              1.85
                           28.4
              1.78
                     89.8 28.3
## 4 Male
## 5 Male
              1.73
                     70.3 23.5
## 6 Female
              1.57
                     85.7 34.8
```

tail(survey)

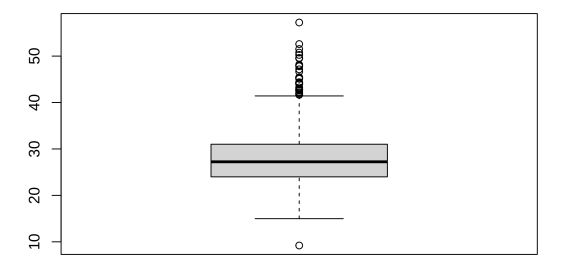
```
## # A tibble: 6 x 4
##
            height weight
     sex
     <fct>
            <dbl> <dbl> <dbl>
## 1 Female
                     95.7 35.2
             1.65
## 2 Male
             1.8
                     79.4 24.5
## 3 Female
             1.73
                     83
                           27.7
## 4 Female
             1.57
                     61.2 24.8
## 5 Male
                     73
                           25.3
              1.7
## 6 Female
             1.55
                     91.2 38.0
```

hist(survey\$bmi)

Histogram of survey\$bmi



boxplot(survey\$bmi)



In the general population, BMI ranges between about 15 to 30. It appears that BMI has been correctly generated in this example. We should investigate the very low and some of the very high values of BMI, but this will be left for another time.

2.7 Summarising data by another variable

We will often want to calculate the same summary statistics by another variable. For example, we might want to calculate summary statistics for BMI for males and females separately. We can do this in in the descriptives function by defining sex as a splitBy variable:

```
library(jmv)
descriptives(data=survey, vars=bmi, splitBy = sex)
```

```
##
    DESCRIPTIVES
##
##
##
    Descriptives
##
##
                              sex
                                         bmi
##
##
      N
                              Male
                                               513
##
                                               627
                              Female
##
      Missing
                              Male
                                                 0
##
                              Female
                                                 0
##
      Mean
                              Male
                                         28.29561
                                         27.81434
##
                              Female
```

```
Median
                                        27.39592
##
                             Male
##
                             Female
                                        26.66667
##
      Standard deviation
                             Male
                                        5.204975
##
                             Female
                                        6.380523
##
      Minimum
                             Male
                                        16.47519
##
                             Female
                                        9.209299
##
                             Male
                                        57.23644
      Maximum
##
                             Female
                                        52.59516
##
```

2.8 Summarising a single column of data

In Module 1, we started with a very simple analysis: reading in six ages, and them using summary() to calculate descriptive statistics. We then went on to use the decriptives() function in the jmv package as more flexible way of calculating descriptive statistics. Let's revisit this analysis:

```
# Author: Timothy Dobbins
# Date: 5 April 2022
# Purpose: My first R script
library(jmv)
age \leftarrow c(20, 25, 23, 29, 21, 27)
# Use "summary" to obtain descriptive statistics
summary(age)
##
      Min. 1st Qu. Median
                              Mean 3rd Ou.
                                               Max.
##
     20.00 21.50
                    24.00 24.17 26.50
                                              29.00
# Use "descriptives" to obtain descriptive statistics
descriptives(age)
```

Error: Argument 'data' must be a data frame

The summary() function has worked correctly, but the descriptives() function has given an error: Error: Argument 'data' must be a data frame. What on earth is going on here? The error gives us a clue here - the descriptives() function requires a data frame for analysis. We have provided the object age: a **vector**. As we saw in Section 1.6.3, a vector is a single column of data, while a data frame is a collection of columns.

In order to summarise a vector using the descriptives() function, we must first convert the vector into a data frame using as.data.frame(). For example:

```
## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 20.00 21.50 24.00 24.17 26.50 29.00
```

```
# Create a new data frame from the vector age:
age_df <- as.data.frame(age)
# Use "descriptives" to obtain descriptive statistics for age_df
descriptives(age_df)</pre>
```

```
##
##
    DESCRIPTIVES
##
    Descriptives
##
##
##
                              age
##
##
                                      6
      Ν
##
      Missing
##
      Mean
                              24.16667
##
      Median
                              24.00000
##
      Standard deviation
                              3.488075
##
      Minimum
                              20.00000
##
      Maximum
                              29.00000
##
```

2.9 Plotting data by another variable

Unfortunately, it is not straight-forward to create separate plots for every level of another variable. We will demonstrate by plotting BMI by sex using our health survey data.

The following steps are not the most efficient way of doing this, but are easy to follow and understand. We first begin by creating two new data frames, for males and females separately, using the subset() function:

```
survey_males <- subset(survey, sex=="Male")
survey_females <- subset(survey, sex=="Female")</pre>
```

Note that we use the **label** for sex, not the underlying numeric value, as sex is a **factor**.

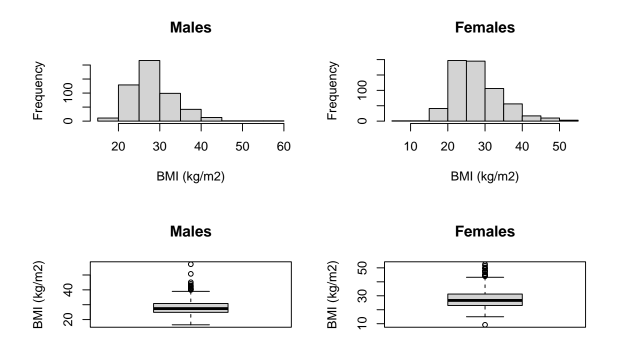
We can now create hisotgrams and boxplots of BMI for males and females separately. To place the graphs next to each other in a single figure, we can use the par function, which sets the *graphics* parameters. Essentially, we want to tell R to split a plot window into a matrix with *nr* rows and *nc* columns, and we fill the cells by rows (mfrow) or columns (mfcols).

For example, to plot four figures in a single plot, filled by rows, we use par(mfrow=c(2,2)). When we are done plotting multiple graphs, we can reset the graphics parameters by submitting par(mfrow=c(1,1)).

```
# Set the graphics parameters to plot 2 rows and 2 columns:
par(mfrow=c(2,2))

# Specify each plot separately
hist(survey_males$bmi, xlab="BMI (kg/m2)", main="Males")
hist(survey_females$bmi, xlab="BMI (kg/m2)", main="Females")

boxplot(survey_males$bmi, ylab="BMI (kg/m2)", main="Males")
boxplot(survey_females$bmi, ylab="BMI (kg/m2)", main="Females")
```



```
# Reset graphics parameters
par(mfrow=c(1,1))
```

2.10 Recoding data

One task that is common in statistical computing is to recode variables. For example, we might want to group some categories of a categorical variable, or to present a continuous variable in a categorical way.

In this example, we can recode BMI into the following categories as suggested by the World Health Organisation [footnote]:

Underweight: BMI < 18.5

• Normal weight: $18.5 \le BMI < 25$

• Pre-obesity: $25 \le BMI < 30$

• Obesity Class I: $30 \le BMI < 35$

Obesity Class II: 35 ≤ BMI < 40

• Obesity Class III: BMI \geq 40

The quickest way to recode a continuous variable into categories is to use the cut command which takes a continuous variable, and "cuts" it into groups based on the specified "cutpoints"

Notice that lower (BMI=0) and upper (BMI=100) bounds have been specified, as both a lower and upper limit must be defined for each group.

If we examine the new bmi_cat variable:

362

##

18

```
summary(survey$bmi_cat)
## (0,18.5] (18.5,25] (25,30] (30,35] (35,40] (40,100]
```

205

97

47

we see that each group has been labelled (a, b]. This notation is equivalent to: greater than a, and less than or equal to b. The cut function excludes the lower limit, but includes the upper limit. Our BMI ranges have been defined to include the lower limit, and exclude the upper limit (for example, greater than or equal to 30 and less than 35).

We can specify this recoding using the right=FALSE option:

411

```
survey$bmi_cat <- cut(survey$bmi,</pre>
                        breaks = c(0, 18.5, 25, 30, 35, 40, 100),
                        right=FALSE)
summary(survey$bmi_cat)
    [0,18.5) [18.5,25)
                           [25,30)
                                      [30, 35)
                                                 [35,40)
##
                                                          [40,100)
##
          18
                    362
                               411
                                           201
                                                      101
                                                                  47
```

Finally, we can specify labels for the groups using the labels option:

```
## Underweight Normal Pre-obesity Obesity Class I
## 18 362 411 201
## Obesity Class II Obesity Class III
## 101 47
```

2.11 Computing binomial probabilities using R

There are two R functions that we can use to calculate probabilities based on the binomial distribution: dbinom and pbinom:

- dbinom(x, size, prob) gives the probability of obtaining x successes from size trials when the probability of a success on one trial is prob;
- pbinom(q, size, prob) gives the probability of obtaining q **or fewer** successes from size trials when the probability of a success on one trial is prob;
- pbinom(q, size, prob, lower.tail=FALSE) gives the probability of obtaining **more than** qsuccesses from size trials when the probability of a success on one trial is prob.

To do the computation for part (a) in Worked Example 2.1, we will use the dbinom function with:

- x is the number of successes, here, the number of smokers (i.e. k=3);
- size is the number of trials (i.e. n=6);
- and prob is probability of drawing a smoker from the population, which is 19.8% (i.e. p=0.198).

Replace each of these with the appropriate number into the formula:

```
dbinom(x=3, size=6, prob=0.198)
```

```
## [1] 0.08008454
```

To calculate the upper tail of probability in part (b), we use the pbinom(lower.tail=FALSE) function. Note that the pbinom(lower.tail=FALSE) function **does not include q**, so to obtain 4 or more successes, we need to enter q=3:

```
pbinom(q=3, size=6, prob=0.198, lower.tail=FALSE)
```

[1] 0.01635325

For the lower tail for part (c), we use the pbinom function:

```
pbinom(q=2, size=6, prob=0.198)
```

[1] 0.9035622

2.12 Computing probabilities from a Normal distribution

We can use the pnorm function to calculate probabilities from a Normal distribution:

- pnorm(q, mean, sd) calculates the probability of observing a value of q or less, from a Normal distribution with a mean of mean and a standard deviation of sd. Note that if mean and sd are not entered, they are assumed to be 0 and 1 respectively (i.e. a standard normal distribution.)
- pnorm(q, mean, sd, lower.tail=FALSE) calculates the probability of observing a value of more than q, from a Normal distribution with a mean of mean and a standard deviation of sd.

To obtain the probability of obtaining 0.5 or greater from a standard normal distribution:

```
pnorm(0.5, lower.tail=FALSE)
```

```
## [1] 0.3085375
```

To calculate the worked example: Assume that the mean diastolic blood pressure for men is 77.9 mmHg, with a standard deviation of 11. What is the probability that a man selected at random will have high blood pressure (i.e. diastolic blood pressure greater than or equal to 90)?

```
pnorm(90, mean=77.9, sd=11, lower.tail=FALSE)
```

[1] 0.1356661

Module 3

Precision: R notes

3.1 Calculating a 95% confidence interval of a mean

3.1.1 Individual data

To demonstrate the computation of the 95% confidence interval of a mean we have used data from Example_1.3.rds which contains the weights of 30 students:

```
library(jmv)
students <- readRDS("data/examples/Example_1.3.rds")
summary(students)

## weight gender
## Min. :60.00 Male :16
## 1st Qu.:67.50 Female:14
## Median :70.00
## Mean :70.00
## 3rd Qu.:74.38
## Max. :80.00</pre>
```

The mean and its 95% confidence interval can be obtained many ways in R. We will use the t.test() function installed in R to calculate the confidence interval:

t.test(students\$weight)

```
##
## One Sample t-test
##
## data: students$weight
## t = 76.029, df = 29, p-value < 2.2e-16
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
## 68.11694 71.88306
## sample estimates:
## mean of x
## 70</pre>
```

The output of the t.test() function gives us the sample mean (70.0 kg) as well as the 95% confidence interval around the mean: 68.1 to 71.9 kg.

Note: the descriptives() function within the jmv package also calculates a 95% confidence interval around the mean. **It is recommended not to use this function** as it currently (as of June 2022) uses a *z* value to calculate the confidence interval, rather than a *t* value.

3.1.2 Summarised data

For Worked Example 3.2 where we are given the sample mean, sample standard deviation and sample size. R does not have a built-in function to calculate a confidence interval from summarised data, but we can write our own.

Note: writing your own functions is beyond the scope of this course. You should copy and paste the code provided to do this.

Module 4

Hypothesis testing

4.1 One sample t-test

Max. :122.00

:35

NA's

We will use data from Example_4.1.rds to demonstrate how a one-sample t-test is conducted in R.

```
bloodpressure <- readRDS("data/examples/Example_4.1.rds")

summary(bloodpressure)

## dbp

## Min. : 24.00

## 1st Qu.: 64.00

## Median : 72.00

## Mean : 72.41

## 3rd Qu.: 80.00
```

To test whether the mean diastolic blood pressure of the population from which the sample was drawn is equal to 71, we can use the t.test command:

```
t.test(bloodpressure$dbp, mu=71)
```

```
##
## One Sample t-test
##
## data: bloodpressure$dbp
## t = 3.0725, df = 732, p-value = 0.002202
## alternative hypothesis: true mean is not equal to 71
## 95 percent confidence interval:
## 71.50732 73.30305
## sample estimates:
## mean of x
## 72.40518
```

The output gives a test statistic, degrees of freedom and a P values from the two-sided test. The mean of the sample is provided, as well as the 95% confidence interval.

Module 5

Comparing two means

5.1 Setting an observation to missing

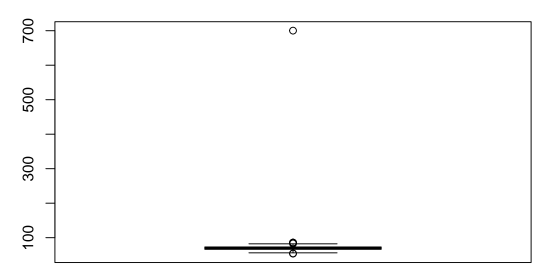
Setting an incorrect observation to missing is straightforward in Stata by using the Data Editor. While RStudio allows browsing a data set as a spreadsheet, it will not let a user replace an observation with a missing value: this should be done using code.

A missing value in R is denoted NA, and this is consistent for any variable type: continuous, string (i.e. character) and even a factor.

Recall the weights data used in Module 2. In viewing a boxplot of weight, we saw an obvious outlier of 700.2kg for ID 58:

```
library(jmv)
sample <- read.csv("data/examples/Weight_s2.csv")
boxplot(sample$weight, xlab="Weight (kg)", main="Boxplot of 1000 weights")</pre>
```

Boxplot of 1000 weights



Weight (kg)

```
subset(sample, weight>200)
## id weight
## 58 58 700.2
```

We previously set this value to 70.2kg using an ifelse() command. Here, let's create a new, cleaned weight variable, and set the incorrect value to missing:

```
sample$weight_clean = ifelse(sample$weight==700.2, NA, sample$weight)
```

Our code will create a new column (called weight_clean) in the sample dataframe. We will test whether weight is equal to 700.2; if this is true, we will assign weight_clean to be NA (i.e. missing), otherwise weight_clean will equal the value of weight.

Let's view the data from ID 58, and summarise the cleaned weight variable using descriptives() and a boxplot:

```
subset(sample, sample$id==58)

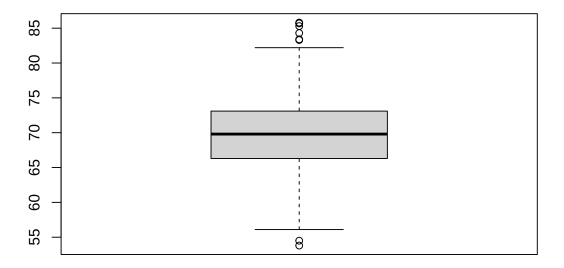
## id weight weight_clean
## 58 58 700.2 NA

descriptives(data=sample, vars=weight_clean)

##
## DESCRIPTIVES
##
## Descriptives
```

```
##
##
                              weight_clean
##
##
      Ν
                                        999
##
      Missing
##
      Mean
                                   69.76406
##
      Median
                                   69.80000
##
      Standard deviation
                                   5.055188
##
      Minimum
                                   53.80000
##
      Maximum
                                   85.80000
##
```

Boxplot of 999 weights



Weight (kg)
(Excluding 1 observation of 700.2kg)

5.2 Checking data for the independent samples t-test

5.2.1 Producing histograms and boxplots by a second variable

We have seen how to create histograms and boxplots separated by a second variable in Module 2 (Section 2.9). We will demonstrate using the birthweight data in Example_5.1.rds.

```
library(jmv)
bwt <- readRDS("data/examples/Example_5.1.rds")
summary(bwt)</pre>
```

```
##
       gender
                  birthweight
##
    Female:56
                 Min.
                         :2.750
    Male :44
                 1st Qu.:3.257
##
##
                 Median :3.450
                         :3.514
##
                 Mean
                 3rd Qu.:3.772
##
##
                 Max.
                         :4.250
```

summary(bwt\$gender)

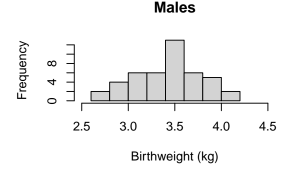
```
## Female Male
## 56 44
```

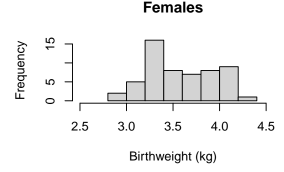
We can create subsets of the birthweight data, subsetted for males and females separately. Note here that gender is a factor, so we need to select based on the factor labels, not the underlying numeric code.

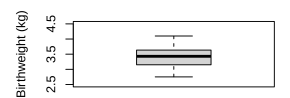
```
bwt_m <- subset(bwt, bwt$gender=="Male")
bwt_f <- subset(bwt, bwt$gender=="Female")</pre>
```

We can now create histograms and boxplots for males and females separately, in the usual way, using the par function to set the graphics parameters to display graphs in a 2-by-2 grid:

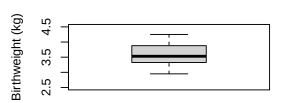
```
\label{eq:par_main} \begin{split} & \mathsf{par}(\mathsf{mfrow} \texttt{=} \mathsf{c}(2,2)) \\ & \mathsf{hist}(\mathsf{bwt} \texttt{\_m\$birthweight}, \ \mathsf{xlim} \texttt{=} \mathsf{c}(2.5,\ 4.5), \ \mathsf{xlab} \texttt{=}"Birthweight} \ (\mathsf{kg})", \ \mathsf{main} \texttt{=}"Males") \\ & \mathsf{hist}(\mathsf{bwt} \texttt{\_f\$birthweight}, \ \mathsf{xlim} \texttt{=} \mathsf{c}(2.5,\ 4.5), \ \mathsf{xlab} \texttt{=}"Birthweight} \ (\mathsf{kg})", \ \mathsf{main} \texttt{=}"Females") \\ & \mathsf{boxplot}(\mathsf{bwt} \texttt{\_m\$birthweight}, \ \mathsf{ylim} \texttt{=} \mathsf{c}(2.5,\ 4.5), \ \mathsf{ylab} \texttt{=}"Birthweight} \ (\mathsf{kg})", \ \mathsf{main} \texttt{=}"Males") \\ & \mathsf{boxplot}(\mathsf{bwt} \texttt{\_f\$birthweight}, \ \mathsf{ylim} \texttt{=} \mathsf{c}(2.5,\ 4.5), \ \mathsf{ylab} \texttt{=}"Birthweight} \ (\mathsf{kg})", \ \mathsf{main} \texttt{=}"Females") \end{split}
```







Males



Females

```
par(mfrow=c(1,1))
```

When we are done plotting multiple graphs, we can reset the plot window by submitting par(mfrow=c(1,1)).

5.2.2 Producing split summary statistics

The descriptives function within the jmv function allows summary statistics to be calculated within subgroups using the splitBy argument:

```
descriptives(data=bwt, vars=birthweight, splitBy=gender)
##
    DESCRIPTIVES
##
##
##
    Descriptives
##
##
                                        birthweight
                              gender
##
##
      Ν
                              Female
                                                  56
##
                              Male
                                                  44
##
      Missing
                              Female
                                                    0
##
                              Male
                                                    0
##
      Mean
                              Female
                                            3.587411
##
                              Male
                                            3.421364
##
      Median
                                            3.530000
                              Female
##
                              Male
                                            3.430000
      Standard deviation
##
                              Female
                                           0.3629788
##
                              Male
                                           0.3536165
##
      Minimum
                              Female
                                            2.950000
##
                              Male
                                            2.750000
##
      Maximum
                              Female
                                            4.250000
##
                              Male
                                            4.100000
```

5.3 Independent samples t-test

##

We can use the ttestIS() (t-test, independent samples) function from the jmv package to perform the independent samples t-test. We include the meanDiff=TRUE and ci=TRUE options to obtain the difference in means, with its 95% confidence interval. We can request a Welch's test (which does not assume equal variances) by the welchs=TRUE option:

```
ttestIS(data=bwt, vars=birthweight, group=gender, meanDiff=TRUE, ci=TRUE)
##
##
    INDEPENDENT SAMPLES T-TEST
##
    Independent Samples T-Test
##
##
                                                        Mean difference
                                                                          SE difference
##
                         Statistic
                                      df
                                               р
                                                                                          Lower
##
##
    birthweight
                   Student's t
                                 2.296556
                                            98.00000
                                                        0.0237731
                                                                        0.1660471
                                                                                      0.07230265
##
```

```
ttestIS(data=bwt, vars=birthweight, group=gender, meanDiff=TRUE, ci=TRUE, welchs=TRUE)
```

```
##
    INDEPENDENT SAMPLES T-TEST
##
##
##
   Independent Samples T-Test
##
##
                         Statistic
                                                         Mean difference
                                                                           SE difference
                                                                                           Lower
                                                                                                       Upp
                                               р
##
##
                                  2.296556
                                             98.00000
                                                        0.0237731
                                                                         0.1660471
                                                                                       0.07230265
                                                                                                     0.022
    birthweight
                   Student's t
               Welch's t
##
                             2.303840
                                        93.54377
                                                   0.0234458
                                                                    0.1660471
                                                                                   0.07207403
                                                                                                0.0229332
##
```

There is no built-in function to calculate an independent t-test from summarised data, nor is there a function within jmv. We can use the tsum.test() function within the BSDA package, with the following syntax:

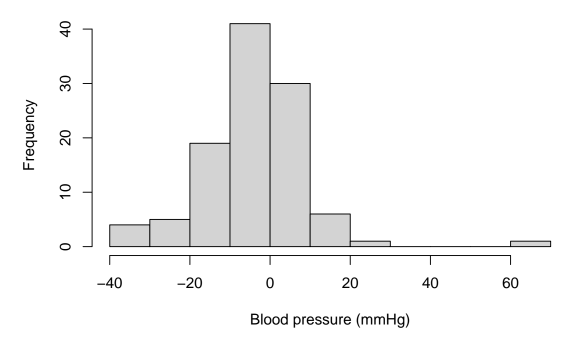
Here we specify the mean, standard deviation and sample size for the first group (on the first line) and the second group (on the second line). We can relax the assumption of equal variances using var.equal=FALSE.

5.4 Checking the assumptions for a Paired t-test

Before performing a paired t-test, you must check that the assumptions for the test have been met. Using the dataset Example_5.2.rds to show that the difference between the pair of measurements between the sites is normally distributed, we first need to compute a new variable of the differences and examine its histogram.

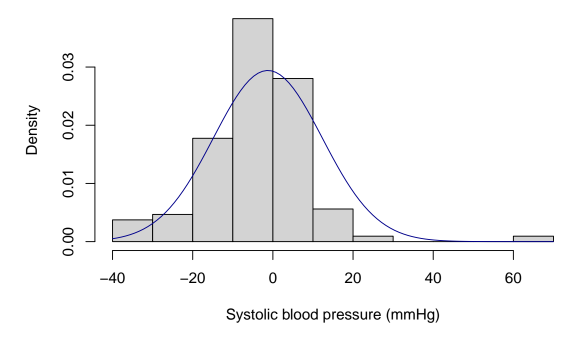
```
sbp <- readRDS("data/examples/Example_5.2.rds")
sbp$diff = sbp$sbp_dp - sbp$sbp_tp
hist(sbp$diff, xlab="Blood pressure (mmHg)", main="Difference in systolic blood pressure")</pre>
```

Difference in systolic blood pressure



We might want to plot a Normal curve over this distribution, as we did in Module 2:





While there is a large difference in blood pressure (around 60 mmHg) that warrents further checking, the curve is roughly symmetric with an approximately Normal distribution.

5.5 Paired t-Test

To perform a paired t-test we will use the dataset Example_5.2.rds. We can perform a paired t-test using the ttestPS() function within the jmv package, where we defined the paired observations as: 'pairs=list(list(i1 = 'variable1', i2 = 'variable2'))

```
ttestPS(data=sbp, pairs=list(list(i1 = 'sbp_dp', i2 = 'sbp_tp')), meanDiff=TRUE, ci=TRUE)
##
##
    PAIRED SAMPLES T-TEST
##
    Paired Samples T-Test
##
##
                                                            Mean difference
                                                                               SE difference
##
                                          df
                             statistic
                                                                                               Lower
##
##
                      Student's t
                                     -0.9621117
                                                  106.0000
                                                             0.3381832
                                                                              -1.261682
                                                                                             1.311368
##
```

The syntax of the ttestPS function is a little cumbersome. The t.test function can be used as an alternative:

```
t.test(sbp$sbp_dp, sbp$sbp_tp, paired=TRUE)

##
## Paired t-test
##
```

5.5. PAIRED T-TEST 65

```
## data: sbp$sbp_dp and sbp$sbp_tp
## t = -0.96211, df = 106, p-value = 0.3382
## alternative hypothesis: true mean difference is not equal to 0
## 95 percent confidence interval:
## -3.861596 1.338232
## sample estimates:
## mean difference
## -1.261682
```

Module 6

Proportions

6.1 95% confidence intervals for proportions

We can use the BinomCI(x=, n=, method=) function within the DescTools package to compute 95% confidence intervals for proportions. Here we specify x: the number of successes, n: the sample size, and optionally, the method (which defaults to Wilson's method).

6.2 Significance test for single proportion

We can use the binom.test function to perform a significance test for a single proportion: binom.test(x=, n=, p=). Here we specify x: the number of successes, n: the sample size, and p: the hypothesised proportion (which defaults to 0.5 if nothing is entered).

```
binom.test(x=54, n=300, p=0.2)

##

## Exact binomial test

##

## data: 54 and 300

## number of successes = 54, number of trials = 300, p-value = 0.4273

## alternative hypothesis: true probability of success is not equal to 0.2

## 95 percent confidence interval:

## 0.1382104 0.2282394

## sample estimates:

## probability of success

## 0.18
```

Active :50

Nausea

Note that the binom.test function also produces a 95% confidence interval around the estimated proportion. This confidence interval is based on the inferior Wald method: the confidence interval derived from the Wilson method is preferred.

6.3 Computing a relative risk and its 95% confidence interval

:19

We will use Worked Example 6.4 to demonstrate calculating a relative risk and its 95% CI:

```
library(jmv)
drug <- readRDS("data/examples/Example_6.4.rds")
summary(drug)

## group side_effect
## Placebo:50 No nausea:81</pre>
```

By using the head() function to view the first six lines of data, we see that both group and side_effect have been entered as factors. Notice the order in which the factor levels are presented: group has the Placebo level defined as the first level, and the Active level defined as the second; side_effect has No nausea defined as the first level, and the Nausea level defined as the second. We will use jmv to calculate relative risks, odds ratios and risk differences. To calculate these estimates correctly, we must define the positive exposure and positive outcome to be the first level of a factor. When defining an exposure for example, we should define the active treatment or the positive exposure as the first category. When defining an outcome, we should define the category of interest (e.g. disease, or side effect) as the first category.

In this example, we will define Active as the first level in the group factor, and Nausea to be the first level of the side_effect factor.

We can do this using the relevel() function, which re-orders the levels of a factor so that the level specified is defined as the first level, and the others are moved down:

```
# Define "Active" as the first level of group:
drug$group <- relevel(drug$group, ref="Active")

# Define "Nausea" as the first level of side_effect:
drug$side_effect <- relevel(drug$side_effect, ref="Nausea")</pre>
```

Upon re-leveling the factors, we can check that the levels of interest have been defined as the first levels:

```
summary(drug)

## group side_effect
## Active :50 Nausea :19
## Placebo:50 No nausea:81
```

To construct the 2-by-2 table and calculate a relative risk, we use the contTables() function in jmv. We request the row-percents using pcRow = TRUE and the relative risk and confidence interval using relRisk = TRUE:

```
##
##
    CONTINGENCY TABLES
##
##
    Contingency Tables
##
##
      group
                                     Nausea
                                                    No nausea
                                                                   Total
##
                   Observed
                                                                           50
##
      Active
                                             15
                                                            35
##
                   % within row
                                      30.00000
                                                     70.00000
                                                                   100.00000
##
##
      Placebo
                   Observed
                                                                           50
                                              4
                                                            46
                   % within row
                                       8.00000
                                                     92.00000
                                                                   100.00000
##
##
      Total
                   Observed
##
                                             19
                                                            81
                                                                          100
                   % within row
##
                                      19.00000
                                                     81.00000
                                                                   100.00000
##
##
##
##
    x<sup>2</sup> Tests
##
##
             Value
                           df
                                  р
##
##
      X<sup>2</sup>
             7.862248
                                  0.0050478
##
      Ν
                   100
##
##
##
##
    Comparative Measures
##
##
                          Value
                                          Lower
                                                       Upper
##
      Relative risk
                          3.750000
##
                                          1.337540
                                                       10.51370
##
##
         Rows compared
```

If you only have the cross-tabulated data (i.e. aggregated), you will need to enter your data into a new data frame. For example, to recreate the above analyses, we can re-write the 2-by-2 table as follows:

Group	side_effect	Number
Active	Nausea	15
Active	No nausea	35
Placebo	Nausea	4
Placebo	No nausea	46

We can enter these data in a dataframe, comprising three vectors, as follows:

```
drug_aggregated <- data.frame(
  group = c("Active", "Active", "Placebo", "Placebo"),
  side_effect = c("Nausea", "No nausea", "No nausea", "No nausea"),
  n = c(15, 35, 4, 46)
)</pre>
```

We need to define group and side_effect as factors. Here we must define the levels in the order we want the categories to appear in the table. Note that as group and side_effect are entered as text

##

Rows compared

variables, we can omit labels command when defining the factors, and the factor will be labelled using the text entry:

```
drug_aggregated$group <- factor(drug_aggregated$group, levels=c("Active", "Placebo"))
drug_aggregated$side_effect <- factor(drug_aggregated$side_effect, levels=c("Nausea", "No nausea"))</pre>
```

We can calculate the relative risk using the summarised data in the same was done previously. However, we need to include the number of observations in each cell using the counts command:

```
contTables(data=drug_aggregated,
           rows=group, cols=side_effect, count=n,
           pcRow=TRUE, relRisk = TRUE)
##
##
    CONTINGENCY TABLES
##
    Contingency Tables
##
##
##
      group
                                   Nausea
                                                 No nausea
                                                               Total
##
##
      Active
                  Observed
                                           15
                                                         35
                                                                       50
                                    30.00000
##
                  % within row
                                                  70.00000
                                                               100.00000
##
##
      Placebo
                  Observed
                                                                       50
                                                         46
##
                  % within row
                                     8.00000
                                                   92.00000
                                                               100.00000
##
##
      Total
                  Observed
                                           19
                                                                      100
##
                  % within row
                                    19.00000
                                                   81.00000
                                                               100.00000
##
##
##
    x² Tests
##
##
##
            Value
                         df
##
            7.862248
                           1
                                0.0050478
##
      χ²
##
                  100
##
##
##
##
    Comparative Measures
##
##
                                                    Upper
                        Value
                                        Lower
##
##
      Relative risk
                        3.750000
                                        1.337540
                                                    10.51370
##
```

6.4 Computing a difference in proportions and its 95% confidence interval

We can use the contTables function to obtain a difference in proportions and its 95% CI, by specifying diffProp=TRUE:

```
contTables(data=drug,
    rows=group, cols=side_effect,
    pcRow=TRUE, diffProp=TRUE)
```

```
##
##
    CONTINGENCY TABLES
##
##
    Contingency Tables
##
##
                                   Nausea
                                                No nausea
                                                              Total
      group
##
##
      Active
                  Observed
                                          15
                                                        35
                                                                      50
                                    30.00000
                  % within row
                                                  70.00000
##
                                                               100.00000
##
##
      Placebo
                  Observed
                                                        46
                                                                      50
                  % within row
                                     8.00000
                                                  92.00000
##
                                                               100.00000
##
##
      Total
                  Observed
                                          19
                                                        81
                                                                     100
##
                  % within row
                                    19.00000
                                                  81.00000
                                                               100.00000
##
##
##
    x² Tests
##
##
##
            Value
                         df
##
      Χ²
            7.862248
                          1
                                0.0050478
##
                  100
##
      N
##
##
##
##
    Comparative Measures
##
##
                                       Value
                                                       Lower
                                                                      Upper
##
##
      Difference in 2 proportions
                                       0.2200000
                                                       0.07238986
                                                                      0.3676101
##
##
        Rows compared
```

6.5 Computing an odds ratio and its 95% confidence interval

We can use the contTables function to obtain an odds ratio and its 95% CI, by specifying odds=TRUE. Here we will use the summarised HPV data from Module 6.

```
hpv <- data.frame(
  hpv = c("HPV +", "HPV +", "HPV -", "HPV -"),
  cancer = c("Case", "Control", "Case", "Control"),
  n = c(57, 14, 43, 186)
)
hpv$cancer <- factor(hpv$cancer, levels=c("Case", "Control"))
hpv$hpv <- factor(hpv$hpv, levels=c("HPV +", "HPV -"))
contTables(data=hpv,</pre>
```

```
rows=hpv, cols=cancer, count=n,
odds = TRUE)
```

```
##
    CONTINGENCY TABLES
##
##
##
    Contingency Tables
##
##
      hpv
                Case
                         Control
                                     Total
##
      HPV +
                  57
                                        71
##
                              14
                                       229
      HPV -
                  43
                             186
##
                             200
                                       300
##
      Total
                 100
##
##
##
##
    x² Tests
##
             Value
##
                          df
                                 p
##
##
      \chi^2
             92.25660
                           1
                                 < .0000001
##
      Ν
                  300
##
##
##
    Comparative Measures
##
##
##
                     Value
                                   Lower
                                                Upper
##
##
      Odds ratio
                     17.61130
                                   8.992580
                                                34.49041
##
```

Note that 95% confidence intervals for the odds ratio based on jmv may differ from those calculated by Stata. At this stage, the method used by jmv to calculate the confidence interval is not documented.

Bibliography

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