MAST20005/MAST90058: Week 2 Lab & R Reference

Goals: (i) Getting started with R and RStudio; (ii) Basic exploratory data analyses; (iii) Basic graphics.

This is much longer than the usual weekly lab sheet because it includes a substantial guide to get you started. You are not expected to absorb it completely in a single lab session. We suggest you skim it upon first reading and ensure you set aside at least 15 minutes to work through the lab exercises at the end in your first lab class, referring back to previous sections when required.

1 Introduction

R is a popular computational tool for statistical applications. An increasing number of researchers in many disciplines, including social and biomedical sciences, choose R for their work. R is both a programming language and a general-purpose computing environment withing which many classical and modern statistical procedures have been implemented.

R is often the vehicle of choice for research in statistical methodology. Its open source nature allows statisticians to contribute new methods in the form of packages which can be downloaded and installed for free at any time. Besides the ordinary packages supplied in the 'base' version of R, hundreds of contributed packages are available through the CRAN website.

The main advantages of R are the fact that it is freely available and that help is readily available online. It is similar to some other high-level programming packages such as MATLAB (which is not free), but more user-friendly than low-level programming languages such as C++ or Fortran. You can use R as it is, but for educational purposes we prefer to use the RStudio interface (also freely available), which has a usefully organized window layout and several extra features which make working with R easier.

2 Getting started

2.1 Install R and RStudio

Both R and RStudio are installed on the machines in your computer labs. If you wish to install R on your own personal computer, go the R Project homepage and follow the instructions. Likewise, to install RStudio, go to the RStudio homepage and follow the instructions.

2.2 RStudio window layout

- Bottom left: console window (also called command window). Here you can type simple commands after the > prompt and R will then execute your command. This is the most important window because this is where R actually does things.
- Top left: editor window (also called script window). Collections of commands (scripts) can be edited and saved. If you do not see this window, you can open it with File > New R script. Just typing a command in the editor window is not enough, it has to get into the command window before R executes the command. If you want to run a line from the script window (or the whole script), you can click Run or press CTRL+ENTER to send it to the command window.
- Top right: workspace / history window. In the workspace window you can see which data and values R has in its memory. You can view and edit the values by clicking on them. The history window shows what has been typed before.

• Bottom right: files / plots / packages / help window. Here you can open files, view plots (also previous plots), install and load packages or use the help function.

2.3 Working directory

Your working directory is the folder on your computer in which you are currently working. When you ask R to open a certain file, it will look in the working directory for it, and when you tell R to save a data file or figure, it will save it in the working directory.

To find out your current working directory, run:

> getwd()

Before you start working, set your working directory to where all your data and script files are or should be stored. You can do that using the **setwd()** function, for example:

```
> setwd("M:/MyStuff/R/")
```

Make sure that the slashes are forward slashes and that you don't forget the apostrophes. R is case-sensitive, so make sure you write capitals where necessary.

2.4 Packages

R can carry out a large number of statistical analyses and procedures. This functionality is organized into a number of 'packages'. With the standard installation, most common packages are installed. To get a list of all installed packages, go to the packages window or type library().

Alternatively, in RStudio you can use the console packages window (bottom right). If the box in front of the package name is ticked, the package is loaded (activated) and can be used. There are many more packages available on the R website. If you want to install and use a package, for example the package called triangle, you should click install packages in the packages window and type 'triangle' or type the following in the command window:

> install.packages("triangle")

To load the package: check the box in front of 'triangle' or type the following in the command window:

> library(triangle)

2.5 A note on menus and commands in RStudio

Many of the tasks we will teach you, such as loading data, can be done either by R commands or by using the RStudio menus. Unfortunately, the RStudio menus often change from one version to another. To avoid confusion, we will generally only show you how to accomplish the task using commands. These will work with any version of RStudio and also if you use plain R (without RStudio). Nevertheless, we still recommend that you use RStudio because of its intuitive and convenient interface.

3 Examples of R commands

3.1 Using R as a calculator

We can warm up by using R as calculator:

```
62 + 5 - 7 * 9 + 15/3 - 2^2 # ^ means "to the power of"
## [1] 5
```

Note that commands after the symbol # are not executed. This can be used to include comments to lines of code.

The [1] is where the output begins, showing the result of your calculation. The number '1' here refers to the fact that it is first number in the output. If there are many numbers in the output and it is too long to fit on one line (e.g. the result is a long vector, see below), numbers in brackets at the start of each line will tell you which position in the output vector is shown at the start of that line.

As with most calculators, R evaluates operations in the following order:

- 1. Power
- 2. Multiplication & division
- 3. Summation & subtraction

For example, the following two commands give different results:

```
2 + 3 / 4<sup>3</sup>

## [1] 2.046875

2 + (3 / 4)<sup>3</sup>

## [1] 2.421875
```

There are many mathematical functions that are already in R, ready to use. For example the exponential function, exp(), the natural logarithm, log(), logarithm in base 2, log2(), the square root, sqrt() and trigonometric functions such as sin() and cos().

```
log(25)  # Logarithm of 25 using natural base e

## [1] 3.218876

sqrt(4)  # Square root of 4

## [1] 2

pi * 4^2  # Area of a circle of radius 4

## [1] 50.26548

a <- pi * 4^2  # Create the variable "a"

a  # Print "a"

## [1] 50.26548
```

The assignment operator, <-, represents an arrow pointing at the object receiving the value of the expression. Instead of <-, some people prefer to use =, which is more standard in other programming languages. Either can be used in R.

To remove all variables from R's memory, run the following:

```
rm(list = ls())
```

You will see that RStudio then empties the workspace window. If you only want to remove the variable a, you can run rm(a).

3.2 Vectors and matrices

R's most basic way of storing data is as a vector (a 'row' of numbers). This is also known as a 1-dimensional array. It can also arrange data into multi-dimensional arrays. The special case of a 2-dimensional array is called a matrix.

To construct a vector named x, use the R command c() (it means "concatenate") as follows:

```
x <- c(4.1, -1.3, 2.5, -0.6, -21.7)
x
## [1] 4.1 -1.3 2.5 -0.6 -21.7
```

The following creates a new vector **z** by combining two vectors:

```
x <- c(1, 3, 9, 10)

x

## [1] 1 3 9 10

y <- c(30, 35, 4)

y

## [1] 30 35 4

z <- c(x, y)

z

## [1] 1 3 9 10 30 35 4
```

R enables us to generate commonly used sequences of numbers. For example 1:30 is the vector c(1, 2, ..., 29, 30).

```
n <- 5

1:n - 1

## [1] 0 1 2 3 4

1:(n - 1)

## [1] 1 2 3 4
```

Comparing the two sequences tells us that the priority of the colon operator ':' is higher than the negation operator '-' (i.e. the colon gets executed first). In general it has higher priority than any other arithmetic operator ('+', '-', '*', '/').

Elements in vectors can be addressed by standard [i] indexing. For example:

```
z[2]
## [1] 3
z[1:4]
## [1] 1 3 9 10
z[c(1, 2, 6)]
## [1] 1 3 35
z[2] <- 2015</pre>
```

In the last line, one of the elements is replaced with a new number.

The function seq() is another useful tool for generating sequences. The first two arguments, if given, specify the beginning and end of the sequence.

```
seq(1, 2, by = 0.1)
## [1] 1.0 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 2.0

seq(1, 2, length.out = 20)
## [1] 1.000000 1.052632 1.105263 1.157895 1.210526 1.263158 1.315789
## [8] 1.368421 1.421053 1.473684 1.526316 1.578947 1.631579 1.684211
## [15] 1.736842 1.789474 1.842105 1.894737 1.947368 2.000000
```

A matrix is can be created from scratch using the function matrix(). For example, to define the matrix

$$A = \begin{pmatrix} 2 & 4 & -1 \\ -1 & 2 & 3 \end{pmatrix}$$

you can use the function $\mathtt{matrix}()$ on a vector where the elements of A are listed column by column:

```
A <- matrix(c(2, -1, 4, 2, -1, 3), nrow = 2) # create a matrix

A

## [,1] [,2] [,3]

## [1,] 2 4 -1

## [2,] -1 2 3
```

Matrix operations are similar to vector operations:

```
A[1, 2] # extract the element in first 1st row and 2nd column

## [1] 4

A[1, ] # extract the first row

## [1] 2 4 -1

mean(A) # sample average for all the elements

## [1] 1.5
```

Elements of a matrix can be addressed in the usual way: [row, column]. When you want to select a whole row, you leave the spot for the column number empty (the other way around for columns of course). The last line shows that many functions also work with matrices as arguments.

3.3 Characters

Character vectors are used frequently in R, for example as plot labels. When needed they can be entered by quotes characters. For example:

```
u1 <- c("male", "female")
u2 <- c("apple", "pear", "kiwi", "orange")
u1

## [1] "male"  "female"

u2

## [1] "apple"  "pear"  "kiwi"  "orange"</pre>
```

A useful function to combine numeric and character vectors is paste(), which takes an arbitrary number of arguments and concatenates them one by one into character strings. The arguments are, by default, separated in the result by a single blank character, but this can be changed using the named parameter sep. For example:

```
labels <- paste(c("X", "Y"), 1:10, sep = "")
labels
## [1] "X1" "Y2" "X3" "Y4" "X5" "Y6" "X7" "Y8" "X9" "Y10"</pre>
```

3.4 Vectorisation

When you ask R to do a calculation with a vector, it will usually repeat the same operation on each element of the vector. For example, the following does elementwise addition:

```
c(1, 2) + c(2, 5)
## [1] 3 7
```

Such operations are called *vectorised* operations. Many functions and operations in R are vectorised, which usually leads to neater and more concise code.

When the vectors are all of the same length, then the operation is well-defined: R simply applies the operation to each element in turn. If you give R vectors of different lengths, then it will 'loop back' on the shorter vectors and start repeating elements from the start. This is called *recycling* the vector. For example:

```
c(1, 2, 3) + c(2, 5)

## Warning in c(1, 2, 3) + c(2, 5): longer object length is not a multiple of shorter
object length
## [1] 3 7 5
```

Often this is not what you want, so R will sometimes print a warning message as it did above. However, one case where it is very handy is when you wish to recycle a single value:

```
1:5 + 3
## [1] 4 5 6 7 8
```

4 Functions

You can write functions to automate calculations or operations. Some functions are standard in R or in one of the packages. Later you will learn how to program your own functions. Important basic functions to know are:

- max() and min(): select the largest and smallest elements of a vector
- range() is a function whose value is a vector of length two: c(min(x), max(x))
- sort() returns a vector of the same size as the original vector with the elements arranged in increasing order
- length() is the number of elements in a vector
- sum() gives the total of the elements in vector
- prod() gives the product of the elements in vector
- dim() gives the dimensions of multi-dimensional array
- mean() gives the sample mean of the vector, which is the same as sum(x) / length(x)

For example, to compute a sample mean:

```
mean(z)
## [1] 300.5714
```

Within the brackets you specify the arguments. Arguments give extra information to the function. In this case, the argument says which vector of numbers to compute the mean for.

The function rnorm() is a standard R function which creates random samples from a normal distribution:

```
z <- rnorm(10) # generate a vector with 10 observations from N(0,1) z

## [1] 0.60004985 0.48977383 0.38102583 1.25330446 -0.57493310  
## [6] -1.83025957 1.65287576 -2.19681254 -0.51336285 -0.03412181
```

Here $\mathtt{rnorm}()$ is the function and the 10 is an argument specifying how many random numbers you want, in this case 10 numbers (typing n=10 instead of just 10 would also work). Note that entering the same commands again produces 10 new random numbers. Instead of typing the same text again, you can also press the upward arrow key ' \uparrow ' to access previous commands.

If you want 10 random numbers out of normal distribution $N(\mu=10,\sigma^2=2^2)$ you can type:

```
x <- rnorm(10, mean = 10, sd = 2)
x
## [1] 6.320776 7.321068 9.567314 10.022107 11.834722 11.038895 7.461893
## [8] 8.106456 7.729224 6.419676</pre>
```

Thus, we see that the same function (rnorm) may have different interfaces and that R uses named arguments (in this case mean and sd). By the way, the spaces around the ',' and '=' symbols do not matter. Comparing this example to the previous one also shows that for the function rnorm() only the first argument (the number 10) is compulsory, and that R gives default values to the others (these are known as optional arguments). RStudio has a nice feature: when you type rnorm(in the command window and press TAB, RStudio will show the possible options.

4.1 Getting help

There is a vast amount of (free) documentation and help available. Some help is automatically installed. Typing in the console window the command:

> help(boxplot)

gives the documentation on the boxplot() function. It gives a description of the function, all of the possible arguments, the values that are used as default for each optional argument, and various other information. Typing

> example(boxplot)

gives some examples of how the function can be used.

5 Scripts

You can store your commands in files called scripts. These have the extension .R, for example myscript.R. You can open an editor window to edit these files by clicking File and New or Open file.... You can run (send to the console window) part of the code by selecting lines and pressing CTRL+ENTER or click Run in the editor window. If you do not select anything, R will run the line your cursor is on. You can always run the whole script with the console command:

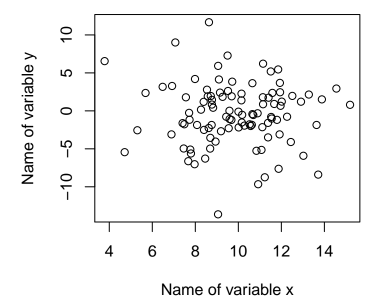
```
> source(myscript.R)
or press CTRL+SHIFT+S.
```

We highly recommend doing and saving all of your work as scripts rather than typing directly into the console window. This makes it easier to correct errors as you go and keep a record of your work.

6 Graphics

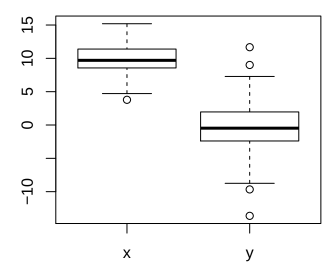
Visualising data is an important component of statistical modelling and data analysis. R can be used to make a large variety of different plots. Here is a simple example:

```
x \leftarrow rnorm(100, mean = 10, sd = 2) # sample of size 100 from N(10, 4)
y \leftarrow rnorm(100, mean = 0, sd = 4) # sample of size 100 from N(0, 16)
plot(x, y, xlab = "Name of variable x", ylab = "Name of variable y")
```



In the first two lines, random numbers are assigned to the variables x and y, which become vectors. In the third line, pairs of values are plotted in the plots window. The following creates a paired boxplot:

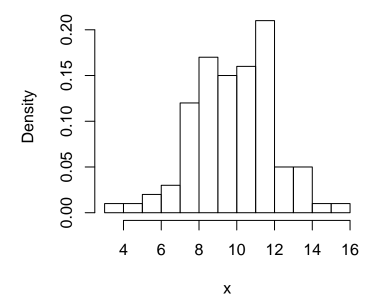
boxplot(x, y, names = c("x", "y")) # creates paired boxplots



Other simple examples are the histogram and normal QQ plots, generated by the simple command:

hist(x, freq = FALSE, nclass = 10)

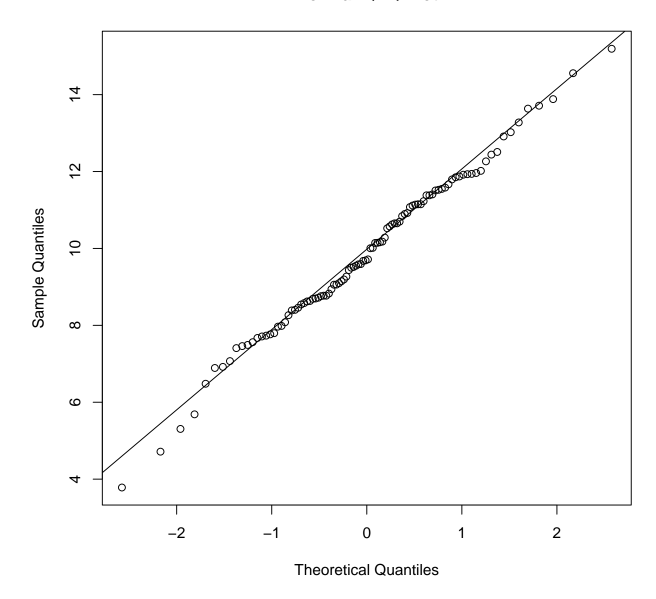
Histogram of x



The second argument specifies that a density histogram is desired (freq = TRUE shows relative frequencies instead), while the third argument specifies the number of bins.

qqnorm(x)
qqline(x)

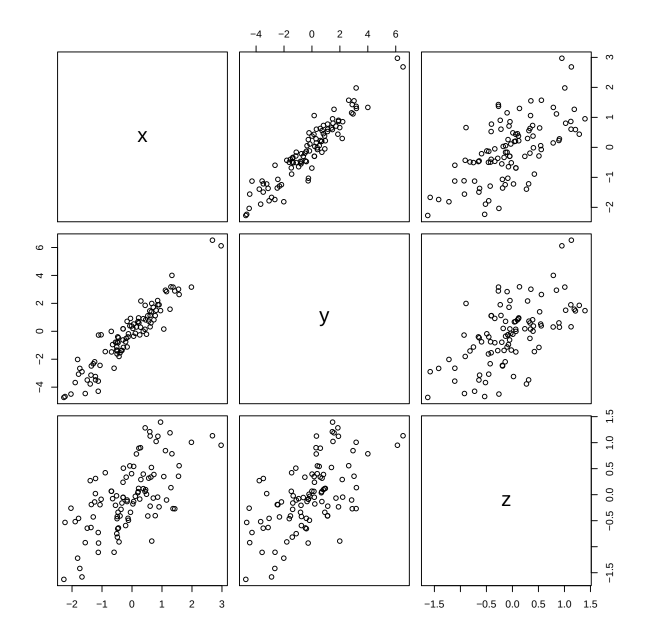
Normal Q-Q Plot



7 Other data structures

Datasets are often organised into data frames, which is like a matrix with names above the columns. This is nice, because you can call and use one of the columns without knowing in which position it is.

```
# generate 3 vectors of length 100
x \leftarrow rnorm(100)
       2 * x + rnorm(100, 0, 0.8)
z \leftarrow 0.5 * x + rnorm(100, 0, 0.5)
t <- data.frame(x, y, z)
                                     # create a data frame
summary(t$x)
                                     # summary statistics for x
##
       Min.
              1st Qu.
                        Median
                                           3rd Qu.
                                    Mean
                                                        Max.
## -2.27169 -0.65598 -0.09537 -0.07988
                                           0.60522
                                                     2.97144
plot(t)
                                     # scatter plot matrix
```



Another basic structure in R is a *list*. The main advantage of lists is that the objects forming the list do not have to be of the same length. (Data frames are actually a special type of list.)

```
L <- list(one = 1, two = c(1, 2), five = seq(0, 1, length.out = 5))
L

## $one
## [1] 1
##

## $two
## [1] 1 2
##

## $five
## [1] 0.00 0.25 0.50 0.75 1.00

L$five + 10

## [1] 10.00 10.25 10.50 10.75 11.00</pre>
```

8 Reading and writing data

There are many ways to write data from within the R environment to files, and also to read data from files. The following lines illustrate some basic steps using console commands:

```
# Construct and store a simple data frame.
t <- data.frame(x = c(1, 2, 3), y = c(30, 20, 10))
t

## x y
## 1 1 30
## 2 2 20
## 3 3 10

write.table(t, file = "mydata.txt", row.names = FALSE) # save file
t2 <- read.table(file = "mydata.txt", header = TRUE) # load file
t2

## x y
## 1 1 30
## 2 2 20
## 3 3 10</pre>
```

The argument row.names = FALSE prevents that row names are written to the fille. Nothing is specified about col.names, so the default option col.names = TRUE is chosen and column names are written to the file. When reading a file note that the column names are also read if header = TRUE. The data frame also appears in the workspace window.

R has several additional packages for reading external data format. For example, the function read.csv() reads CSV formatted text files. Another example is the package foreign which contains functions such as read.dta() & write.dta(), and read.spss() & write.spss(), for dealing with Stata and SPSS formats respectively.

9 Missing data

When you work with real data, you will encounter missing values. When a data point is not available, you write NA instead of a number.

```
x <- c(rnorm(10), NA, rnorm(2))
```

Computing statistics of incomplete data sets is usually an undefined operation, and the output itself will also be NA. For example, R will say that it doesn't know what the smallest value of \mathbf{x} is:

```
min(x)
## [1] NA
```

If you do not care about missing data and want to compute the statistics with the values that are present, use the additional argument na.rm = TRUE:

```
min(x, na.rm = TRUE)
## [1] -2.191828

mean(x, na.rm = TRUE)
## [1] -0.1324518
```

10 Conditional execution and loops

In R, commands can be grouped together by braces, { expression1; expression2; ... }, and the result of the group expression is the result of the last expression being evaluated. We can use single expressions or groups of expression to construct conditional statements with the following syntax:

```
> if (expression1) expression2 else expression3
```

where expression1 is a condition resulting in a logical value TRUE or FALSE. If expression1 is true, then expression2 is evaluated. If expression1 is false, then expression3 is evaluated instead.

Let us consider a sample of n = 10 observations from a normal N(0, 1),

```
x <- rnorm(10)
```

The following code checks whether the mean is greater than the median and prints a string with the outcome.

```
if (mean(x) > median(x)) {
    "The mean is greater than the median"
} else {
    "The mean is smaller than the median"
}
## [1] "The mean is greater than the median"
```

There is also a vectorised version of the if-else statement, in the form a function called ifelse(). The syntax for it is ifelse(condition, a, b).

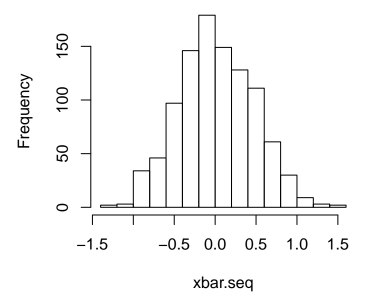
Suppose we want to carry out a task a fixed number of times, say B. To this end, it is convenient to construct for-loops. In a for-loop, you specify a list of possible values and also the name of a variable, which will take each of those values in turn and perform a calculation.

> for (name in expression1) expression2

where name is the loop variable and expression1 is a vector expression (in many cases a sequence like 1:100). From example, consider the drawing B = 1000 samples of size n = 5 from a N(0, 1). For each sample, we compute \hat{x} , and store the results in a vector of length B:

```
B <- 1000  # number of runs
n <- 5  # sample size
xbar.seq <- 1:B  # a vector of size to be filled with means
for (i in 1:B) {
    sample <- rnorm(5)
    xbar.seq[i] <- mean(sample)
}
hist(xbar.seq)  # plot the results</pre>
```

Histogram of xbar.seq



11 Write your own functions

The R language allows the user to create new functions. These are true R functions that are stored in a special internal form and may be used in further expressions and so on. In the process, the language gains enormously in power, convenience and elegance, and learning to write useful functions is one of the main ways to make your use of R comfortable and productive.

A function is defined by an assignment of the form:

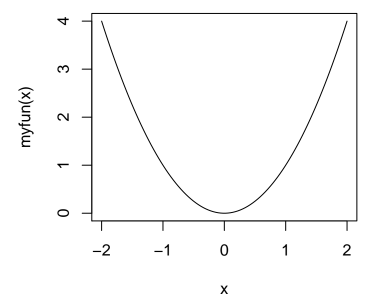
```
> namefunction <- function(argument_1, argument_2, ...) { expression }</pre>
```

For example

```
myfun <- function(x) {  # Specifies function name and argument
    y <- x^2  # Specifies what the function should do
    return(y)  # Returned value
}
myfun(1.5)  # Computes 1.5^2

## [1] 2.25

x <- seq(-2, 2, length.out = 100) # Plots the new function
plot(x, myfun(x), type = "l")</pre>
```



Next write a function to compute the median. A simple algorithm is to sort the data and take the middle element:

```
mymedian <- function(x) {
    n <- length(x)
    m <- sort(x)[(n + 1) / 2]
    return(m)
}</pre>
```

Note that this function operates without an error message for n even or odd, but does not return the correct result for even n. For even sample sizes, we should take the average of the two middle values:

```
mymedian <- function(x) {
    n <- length(x)
    if (n %% 2 == 1) { # odd
        med <- sort(x)[(n + 1) / 2]</pre>
```

Our new function seems to operate correctly and gives a results matching the default median() function.

Exercises

- 1. Calculate $\sum_{i=1}^{100} \ln(i)$.
- 2. Let $X \sim N(1,2)$. What is $\mathbb{E}(X^2)$? Approximate this by simulating a large number of normal random variables and doing an appropriate calculation.
- 3. Use the help system to find out what the qnorm() function does. Explain the result of running qnorm(0.1). What about qnorm(0.1, lower.tail = FALSE)? For what value of a will the command qnorm(a) return the same value as the previous one?
- 4. Write a function explpdf() that calculates the pdf of an exponential distribution with mean 1. Compare the output of your function against the in-built function that does the same calculation (dexp()). Remember to check it works for all inputs: for example, what is the correct value of explpdf(-1)?
- 5. Do question 6(c) from the tutorial problems.
- 6. Do question 7(b) from the tutorial problems.
- 7. Do question 2(c) from the tutorial problems by simulation (rather than using the Central Limit Theorem).