Introduction to R

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- 1. Getting started & R basics
 - Downloading and installing R
 - Organization of RStudio
 - Object types, comments, etc
- 2. Tidyverse
 - Installing and loading packages
 - Importing, tidying, processing & visualizing data.
- 3. Working with matrices in R
 - Creating vectors & matrices.
 - Matrix multiplication, inverse, determinant etc.
 - Application: OLS estimator
- 4. Functions, control structures & loops in R
 - User written functions, if-else structures, for & while loops.
 - Application: maximum likelihood estimation

1. Getting started & R basics

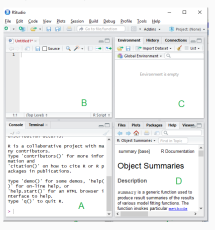
Getting started

- Download this slide deck, example datasets and exercises from https://github.com/hhsievertsen/R_introduction
- 2. Download R from stats.bris.ac.uk/R/ and install it.
- 3. Download RStudio https://rstudio.com/ and install it.
- 4. Open R Studio

For more on how to install R and RStudio see chapter 3 in "R Programming for Data Science" (Peng, 2019)

Organisation of RStudio

- A. Console
- B. Script editor
- C. Overview of objects
- D. Documentation/plots/file browser/packages



4

R basics: R as a calculator

R as a calculator

• We can use R as a calculator. Try typing the following in console and press enter:

5+3

[1] 8

- You can also type 5+3 in the script editor, highlight what you just wrote and click Ctrl+Enter.
- Using the script editor, the keyboard combination Ctrl+Enter executes the current line or the selected area.

R basics: the assignment operator

The assignment operator: <-

value1<-5

- The number five is assigned to an object named "value1".
- We can also achieve this using = instead of <- , but I recommend getting used to using <- as it will become of advantage later on.
- We can also use named objects in the calculator approach:

value1<-5 value1+3

[1] 8

R basics: print()

Printing

We can ask to display the content of an object using print()

```
value1<-5
value2<-3
value3<-value1+value2
print(value3)</pre>
```

```
## [1] 8
```

R returns the value of an expression automatically, this is called automatic printing.

```
value1<-5
value2<-3
value3<-value1+value2
value3
```

```
## [1] 8
```

Automatic printing is disabled in loops, functions etc (more on that later).

R basics: functions

R functions

- print() is an example of a R function.
- The name of this function is print
- The function options (called arguments) go inside the ().
- This is general R syntax:
- If you include () after a name, R knows it is a function. If you don't include () R knows it is not a function.
- Functions can accept many arguments inside the ().
- Ordered arguments

print(value3,TRUE)

• Named arguments

print(x=value3,quote=TRUE)

8

R basics: object type

Object types

```
var<-TRUE
typeof(var)
## [1] "logical"
var<-41.
typeof(var)
## [1] "integer"
var<-4141.2
typeof(var)
## [1] "double"
var<-"Hello1"
typeof(var)
## [1] "character"
```

• Additional types: NULL,raw, complex, list, expression

Vectors

• We **combine** several objects in a vector using the c() function.

```
value1<-5
value2<-3
value3<-value1+value2
vector1<-c(value1,value2,value3)
print(vector1)</pre>
```

```
## [1] 5 3 8
```

a list is homogeneous: all objects are coerced to be of the same type.

```
object1<-414.041
object2<-"hello!"
vector2<-c(object1,object2)
print(vector2)</pre>
```

```
## [1] "414.041" "hello!"
(all objects are strings)
```

R basics: entering values

A range from 1 to 17:

vector < -seq(1,17,by=0.5)

```
vector<-1:17
print(vector)</pre>
```

- ## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17
 - A range from 1 to 17 in steps of 0.5 using seq():

```
print(vector)
## [1] 1.0 1.5 2.0 2.5 3.0 3.5 4.0 4.5 5.0 5.5 6.0 6.5 7.0 7.5
```

A vector of length 12 with missing values (NA) using rep():

```
vector<-rep(NA,12)
print(vector)</pre>
```

- ## [1] NA NA
 - A vector of length 12 with draws from a normal distribution:

```
vector<-rnorm(mean=0,sd=1,n=12)
print(vector)</pre>
```

R basics: comments

Comments

- We should annotate our R script with comments about what we are doing.
- Problem: R will try to execute or comments as R code.
- Solution: Content after the # symbol is ignored by R.

```
# This is line is ignored by R
this is not ignored by R # but this is
```

R basics: useful functions

Create a list (can be heterogenous) using list()

```
a<-"Hello"
b<-1:7
c<-list(a,b)
c

## [[1]]
## [1] "Hello"
##
## [[2]]
## [1] 1 2 3 4 5 6 7</pre>
```

• Get the length of an object using length():

```
vector<-seq(0,3,by=0.02)
obj<-length(vector)</pre>
```

• Concatenate strings with paste():

[1] "Hello Bristol!"

```
obj1<-"Hello"
obj2<-"Bristol"
obj3<-paste(obj1,obj2,"!",sep=" ")
print(obj3)</pre>
```

R basics: working directory and documentation

Working directory

We specify the working directory with 'setwd()'.
 (the default location for saving and loading files.)

setwd("C:\\Users\\hs17922\\Documents")

■ Note: use \\ instead of \.

R documention

- Most functions in R are well documented.
- We can access the documentation by typing ?nameoffunction. For example:

?setwd

Getting started and R basics - summary

- We assign objects using the the assignment operator <-.
- We specify working directory with setwd().
- We access R documentation for the function called functionname with ?functionname.
- We use the # to add comments to our script.
- R has five atomic classes of objects: character, numeric, integer, integer, complex (not covered) and logical.
- We create vectors with c() and lists with list().
- Entering values using rep() and seq().
- Lists are homogeneous and only contain one object class.

For more details see chapter 4 in "R Programming for Data Science" (Peng, 2019)

2. Tidyverse

Installing and loading packages

- R is powerful.
- R with extra packages is very powerful.
- tidyverse is a collection of packages (ggplot2, tidyr,readr, dplyr and more) that are
 useful for working with data.
- to install a package (examplified by "tidyverse").

(We need to do this only once on every system.)

install.packages("tidyverse")

to load a packages (examplified with "tidyverse").

(We have to do that once for every R session.)

library("tidyverse")

Loading data into R

Data formats

- Datasets come in many formats depending on how they were created and saved (Excel, Stata, etc).
- R can load many types of datasets (but sometimes we have to load a special package to load a specific format).
- read_csv() from the readr package (included in tidyverse) is convenient for loading datasets ending on ".csv".
- Note that read.csv() is a slightly different function.

Loading data with read_csv()

```
mydataset<-read_csv("example_data1.csv")</pre>
```

```
##
## cols(
    person_id = col_character(),
    school_id = col_double(),
    summercamp = col_double(),
##
    female = col double().
##
##
    parental_schooling = col_double(),
    parental_lincome = col_double(),
    test_year_1 = col_double(),
##
    test_year_2 = col_double(),
##
    test_year_3 = col_double(),
    test_year_4 = col_double(),
    test_year_5 = col_double(),
##
    test_year_6 = col_double(),
##
    test_year_7 = col_double(),
##
    test_year_8 = col_double(),
##
    test year 9 = col double()
## )
```

- The dataset named "example_data1.csv" in the current working directory is now loaded in R under the name mydataset.
- The dataset is loaded with 15 columns.
- The first variable is a character (i.e. text) type variable and all outhers are double precision floating point numbers.

For more details on importing data see chapter 11 in "R for data science: import, tidy,

Viewing data

What is in mydataset?

- print() will (attempt to) display the full dataset in the console. Not feasible for large datasets.
- head() displays the first six observations in the dataset.

head(mydataset)

```
## # A tibble: 6 x 15
    person_id school_id summercamp female parental_schooling parental_lincome test_year_1 test_year_2 test_year_3 test_year_4
##
    <chr>>
                   <db1>
                               <dbl> <dbl>
                                                         <db1>
                                                                           <db1>
                                                                                       <db1>
                                                                                                    <db1>
                                                                                                                 <db1>
                                                                                                                             <dbl>
## 1 p1
                       5
                                   1
                                          Ω
                                                             14
                                                                            15.3
                                                                                       3.25
                                                                                                     2.99
                                                                                                                 2.58
                                                                                                                             2.16
## 2 p2
                      14
                                                             11
                                                                            14.0
                                                                                       0.993
                                                                                                     1.59
                                                                                                                 1.16
                                                                                                                             0.817
## 3 p3
                       7
                                          0
                                                             13
                                                                            15.1
                                                                                       1.82
                                                                                                     1.02
                                                                                                                 2.38
                                                                                                                             1.88
## 4 p4
                       8
                                          0
                                                             14
                                                                            15.3
                                                                                       2.15
                                                                                                     2.99
                                                                                                                 2.01
                                                                                                                             2.23
## 5 p5
                                          1
                                                             14
                                                                            15.7
                                                                                       3.03
                                                                                                     3.17
                                                                                                                 2.66
                                                                                                                             3.02
## 6 p6
                      26
                                          0
                                                            12
                                                                            14.0
                                                                                       1.52
                                                                                                     1.55
                                                                                                                             1.53
```

- tail() displays the last six observations in the dataset.
- We can specify the number of rows to show with n=6: head(mydataset,n=6) or tail(mydataset,n=6) (to show 6 rows, the default).
- View() opens the dataset in a viewer.

Tidying data I

Tidy data

- The tidy data principles state that each variable must have its own column and each observation must have its own row.
- The example dataset is not tidy.
 - The variables test_year_1 to test_year_9 violate the tidy data principles.
 - The variables contain information about test scores.
 - The values 1,2,..., 9, and 9 are information about the year of the test score, this should be stored in rows for a variable called year.
- The function pivot_longer() (from the tidyr package) gathers several columns in one column (makes the dataset longer).
- The function pivot_wider() (from the tidyr package) spreads one column to several columns (makes the dataset wider).

Tidying data II

pivot_longer()

- pivot_longer(data, cols, names_to, values_to=)
- data: the name of the dataset.
- cols: the columns to convert.
- names_to: the new variable where the information that is currently in the column headers (for example the test year) should be stored.
- values_to: the variable where the values from the old rows are to be stored.

```
## # A tibble: 6 x 8
## person_id school_id summercamp female parental_schooling parental_lincome year test_score
                <db1>
                          <db1> <db1>
                                                <db1>
                                                               <dbl> <chr>
## <chr>
                                                                                   <db1>
## 1 p1
                                                               15.3 test_year_1
                   5
                                                   14
                                                                                   3.25
## 2 p1
                                                               15.3 test_year_2 2.99
                                                   14
                                                               15.3 test_year_3
## 3 p1
                   5
                                                   14
                                                                                   2.58
## 4 p1
                             1 0
                                                   14
                                                               15.3 test_year_4
                                                                                   2.16
## 5 p1
                             1
                                   Ω
                                                   14
                                                               15.3 test_year_5
                                                                                   2.61
## 6 p1
                                   0
                                                   14
                                                               15.3 test_year_6
                                                                                   3.10
```

(I used options(dplyr.width = Inf) to specify the number of columns to print.)

For more details on pivot_longer() data see section 12.3.1 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Tidying data III

6 p6

pivot_wider()

- pivot_wider(data, names_from, values_from=)
- data: the name of the dataset
- names from: the new columns get their names from this variable.
- values_from: the new columns get their values from this variable.

```
dirtydata<-pivot wider(tidydata.
                       names from="year", values from="test score")
options(dplyr.width = Inf)
head(dirtydata)
```

```
## # A tibble: 6 x 15
    person_id school_id summercamp female parental_schooling parental_lincome test_year_1 test_year_2 test_year_3 test_year_4
   <chr>
                   <db1>
                             <dbl> <dbl>
                                                                                    <dbl>
                                                                                                <dbl>
                                                                                                            <db1>
                                                       <db1>
                                                                        <dbl>
                                                                                                                        <dbl>
## 1 p1
                                                                                                 2.99
                      5
                                 1
                                                          14
                                                                         15.3
                                                                                    3.25
                                                                                                             2.58
                                                                                                                        2.16
## 2 p2
                     14
                                        1
                                                          11
                                                                         14.0
                                                                                    0.993
                                                                                                 1.59
                                                                                                             1 16
                                                                                                                        0.817
## 3 p3
                      7
                                        0
                                                          13
                                                                         15.1
                                                                                    1.82
                                                                                                1.02
                                                                                                             2.38
                                                                                                                        1.88
## 4 p4
                      8
                                 1
                                        0
                                                          14
                                                                         15.3
                                                                                    2.15
                                                                                                 2.99
                                                                                                             2.01
                                                                                                                        2.23
## 5 p5
                                 1
                                        1
                                                          14
                                                                         15.7
                                                                                    3.03
                                                                                                 3.17
                                                                                                             2.66
                                                                                                                        3.02
```

12

14.0

1.52

1.55

1.10

0 (I used options(dplyr.width = Inf) to specify the number of columns to print.)

26

0

For more details on pivot_wider() data see section 12.3.2 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016). 1.53

Data processing I

filter

- filter(data, criteria)
- We filter specific rows (observations) of a dataset using the filter() function (from the dplyr package).
- data: the name of the dataset.
- ...: the filtering criteria.

```
filtered_data<-filter(tidydata,year=="test_year_2")
options(dplyr.width = Inf)
head(filtered_data)</pre>
```

```
## # A tibble: 6 x 8
   person_id school_id summercamp female parental_schooling parental_lincome year
                                                                                          test score
##
   <chr>>
                   <dh1>
                              <db1> <db1>
                                                        <14h1>
                                                                        <dbl> <chr>
                                                                                                <14h>>
## 1 p1
                      5
                                                           14
                                                                         15.3 test year 2
                                                                                                2 99
                                         1
                                                                         14.0 test_year_2
## 2 p2
                      14
                                                          11
                                                                                                1.59
                                        0
                                                                         15.1 test_year_2
                                                                                                1.02
## 3 p3
                                                          13
## 4 p4
                                        0
                                                          14
                                                                         15.3 test_year_2
                                                                                                2.99
                                                                         15.7 test year 2
## 5 p5
                      9
                                         1
                                                           14
                                                                                                3.17
## 6 p6
                      26
                                                          12
                                                                         14.0 test year 2
                                                                                                1 55
```

(I used options(dplyr.width = Inf) to specify the number of columns to print.)

For more details on filter() see section 5.2 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing II

select

- select(data, ...)
- We select specific columns (variables) of a dataset using the select() function (from the dplyr package).
- data: the name of the dataset.

(I used options(dplyr.width = Inf) to specify the number of columns to print.)

...: the name (or number) of the columns to keep. To remove a variable, add a
 "-" in front of the variable.

```
selected_data<-select(filtered data.
                         c(person_id,summercamp,test_score))
options(dplyr.width = Inf)
head(selected data)
## # A tibble: 6 x 3
  person_id summercamp test_score
## <chr> <dbl>
                      <dbl>
                 1 2.99
## 1 p1
                 0 1.59
## 2 p2
## 3 p3
                1
                      1.02
## 4 p4
                       2.99
## 5 p5
                 1
                       3.17
## 6 p6
                 0
                      1.55
```

For more details on select() see section 5.4 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing III

rename

- rename(data, newname1=oldname1, newname=oldname,...)
- We rename columns using the rename function
- data: the name of the dataset.
- newname1: the new name of the first column to rename.
- oldname1: the old name of the first column to rename.
- newname2: the new name of the second column to rename.
- ...

(I used options(dplyr.width = Inf) to specify the number of columns to print.)

For more details on rename() see section 5.4 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing IV

mutate

- mutate(data, nameofnewvariable=expression, ...)
- We create and modify columns using the mutate function
- data: the name of the dataset.
- nameofnewvariable: the name of the first column to rename.
- expression: the definition of the new variable
- . . .

```
## # A tibble: 6 x 5
  person_id camp score camptest constant
## <chr> <dbl> <dbl>
                    <db1>
                          <db1>
## 1 p1 1 2.99 2.99
## 2 p2 0 1.59 0
## 3 p3 1 1.02 1.02
         1 2.99 2.99
## 4 p4
           1 3.17 3.17
## 5 p5
                             1
## 6 p6
       0 1.55 0
                             1
```

(I used options(dplyr.width = Inf) to specify the number of columns to print.)

For more details on mutate() see section 5.5 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing V

merge

- merge(x,y,by="matchingvar")
- We merge two datasets using the merge() function.
- x: the name of the first dataset.
- y: the name of the second dataset.
- matchingvar: rows with the same value of matchingvar in both x and y are matched.
-

```
myotherdataset<-read_csv("example_data2.csv")
merged_data<-merge(mutated_data,myotherdataset,by="person_id")
options(dplyr.width = Inf)
head(merged_data)</pre>
```

(I used options(dplyr.width = Inf) to specify the number of columns to print. Some output is hidden.)

For more details on merge() see section 13.4.7 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing VI

The pipe: %>%

- Throw the left-hand side value forward into the right-hand side expression.
- So f(x) can be written as as x%>%f().
- Example

```
renamed_data<-rename(mydataset,score=test_score)
mutated_data<-mutate(renamed_data,camptest=score*camp)
filtered_data<-filter(mutated_data,female==1)
selected_data<-select(filtered_data,person_id,camptest)</pre>
```

can be written as:

For more details on the pipe see chapter 18 in "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Data processing VII

group_by & summarise

- group_by(data,...)
 - data: the name of the dataset.
 - ...: the names of the columns to group the dataset by.
- summarise(data, varname=expression)
 - summarises the dataset on the group_by (if defined) level.
 - data: the name of the dataset.
 - varname: the name of the new variable.
 - expression: the definition of the new variable.

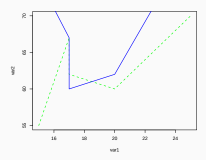
For more details see "section 5.6 in R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

Base graphics I

For more details see R Base Graphics: An Idiot's Guide.

30

Base graphics II



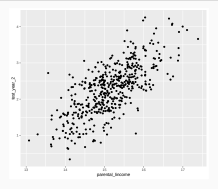
ggplot2 a grammar for graphics

- ggplot(data,aes(x,y,...))
 - data: the name of the dataset.
 - ...: aes() the aesthetic mappings.
 - x the variable to plot on the x-axis.
 - y the variable to plot on the y-axis.
- +geom_line()
 - + add a layer to the ggplot object
 - geom_line() add a line chart using the data and the aesthetic mappings specified in ggplot() (geom_line inherits the settings specified in ggplot()).
- +geom_point(data,aes(x,y,...))
- + add a layer to the ggplot object
- geom_point() add a scatter plot using the data and the aesthetic mappings specified specified within geom_point().
- +theme(): specify theme settings (colors, position etc.)
- +labs(): specify axes titles, chart title, caption, legend titles, etc..

For more details see "chapter 3 in R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

ggplot2 a grammar for graphics

```
ggplot(mydataset,aes(x=parental_lincome,y=test_year_2))+
geom_line()
```



Tidyverse summary

- Tidyverse package tools for working with data: load, tidy, process, visualize data.
- We install packages with install.package().
- We load packages with library().
- We use pivot_wider() and pivot_longer() to tidy the dataset.
- We use mutate() to create new/modify columns in our dataset.
- We use select() to specify which columns to keep/remove.
- We use filter() to specify which rows to keep.
- We use group_by() and summarise() to create aggregate statistics.
- We use ggplot() to create charts.
- We use merge() to merge datasets.

For more details on tidyverse see "R for data science: import, tidy, transform, visualize, and model data." (Wickham & Grolemund, 2016).

3. Working with matrices in R

A vector

• We would like to enter the following vector A into R

$$A = \begin{bmatrix} 3 \\ 5 \\ 4 \end{bmatrix}$$

• We already know how to do this. We simply use c():

$$A < -c(3,5,4)$$
A

[1] 3 5 4

• Note that R prints it as row, but it is a column vector.

A vector transposed

• To verify that A is really a column vector, let's consider the transpose of A:

$$A^T = [3, 5, 4]$$

- which we can obtain in R by means of the transpose function, t():

t(A)

```
## [,1] [,2] [,3]
## [1,] 3 5 4
```

- Where the m in [m,n] refers to the row and the n to the column.
- and let's consider the transpose of the transposed vector to get back to the original A vector.:

t(t(A))

```
## [,1]
## [1,] 3
## [2,] 5
## [3,] 4
```

The matrix function

- We can also explictly create vectors and matrices using the matrix() function.
- To create our A vector, we write:

```
A<-matrix(c(3,5,4),ncol=1)
A
```

```
## [,1]
## [1,] 3
## [2,] 5
## [3,] 4
```

Note the difference between

```
A<-matrix(c(3,5,4),ncol=1)
class(A)
```

```
## [1] "matrix" "array"
```

and

```
A<-c(3,5,4)
class(A)
```

```
## [1] "numeric"
```

Entering a matrix with the matrix function

Creating a 2x2 matrix

• Let us now consider a 2 by 2 matrix *B*:

$$B = \begin{bmatrix} 3, 5 \\ 11, 2 \end{bmatrix}$$

- which we can enter as:

```
B<-matrix(c(3,11,5,2),ncol=2)
B</pre>
```

```
## [,1] [,2]
## [1,] 3 5
## [2,] 11 2
```

- R first fills the first column, then the second etc. . .
- And the transpose of B:

t(B)

```
## [,1] [,2]
## [1,] 3 11
## [2,] 5 2
```

Subsetting

We can extract a subset of an object using squared brackets [row,col]

```
B<-matrix(c(3,11,5,2),ncol=2)
C<-B[2,1]
print(C)</pre>
```

[1] 11

- The first number in the brackets, 2, tells R that we want the second row.
- The second number in the brackets, 1, tells R that we want the first column.
- We therefore extract the value in the second row and the first column.
- We can also extract a range, say that we want the elements in row 1 to 2 and column 1:

```
B<-matrix(c(3,11,5,2),ncol=2)
C<-B[1:2,1]
print(C)</pre>
```

```
## [1] 3 11
```

For more details on subsetting R objects see chapter 9 in "R Programming for Data Science" (Peng, 2019)

Addition

Adding a number to a matrix

• Let α be a number (a scalar), then $\alpha + B$ is:

$$\alpha + B = \begin{bmatrix} \alpha + 3, \alpha + 5 \\ \alpha + 11, \alpha + 2 \end{bmatrix}$$

and in R:

```
B<-matrix(c(3,11,5,2),ncol=2)
alpha<-0.5
C<-alpha+B
print(C)</pre>
```

```
## [,1] [,2]
## [1,] 3.5 5.5
## [2,] 11.5 2.5
```

Adding two matrices together

• Consider the following two matrices

$$C = \begin{bmatrix} 1,2\\3,4 \end{bmatrix} \text{ and } D = \begin{bmatrix} 5,6\\7,8 \end{bmatrix}$$

The sum of these two matrices is then given by::

$$C + D = \begin{bmatrix} 1 + 5, 2 + 6 \\ 3 + 7, 4 + 8 \end{bmatrix}$$

- and in R:

```
C<-matrix(c(1,3,2,4),ncol=2)
D<-matrix(c(5,7,6,8),ncol=2)
E<-C+D
print(E)</pre>
```

```
## [,1] [,2]
## [1,] 6 8
## [2,] 10 12
```

41

Warning: pay attention to dimensions of matrices

• Note: the dimensions have to align when adding two matrices together:

```
E<-matrix(c(1,3,2,4),ncol=1)
F<-matrix(c(5,7,6,8),ncol=2)
E+F</pre>
```

Error in E + F: non-conformable arrays

- E is a 1 × 4 matrix.
- F is a 2×2 matrix.
 - We want to add the element in the first row and first column of E to the element in the first row and the first column of F That's okay.

. . .

 We want to add the element in the third row and first column of E to the element in the third row and the first column of F. That's not doable, because F only has two rows.

Multiplying a matrix with a number

• Let α be a number (a scalar), then αB is:

$$\alpha B = \begin{bmatrix} \alpha \times 3, \alpha \times 5 \\ \alpha \times 11, \alpha \times 2 \end{bmatrix}$$

and in R:

```
B=matrix(c(3,11,5,2),ncol=2)
alpha=0.5
alpha*B
```

```
## [,1] [,2]
## [1,] 1.5 2.5
## [2,] 5.5 1.0
```

Element-wise multiplication

Consider the following two matrices

$$C = \begin{bmatrix} 1, 2 \\ 3, 4 \end{bmatrix} \text{ and } D = \begin{bmatrix} 5, 6 \\ 7, 8 \end{bmatrix}$$

 The Hadamard product (or element-wise multiplication) of matrices C and D is then given by:

$$E = C \circ D = \begin{bmatrix} 1 \times 5, 2 \times 6 \\ 3 \times 7, 4 \times 8 \end{bmatrix}$$

and in R:

```
C=matrix(c(1,3,2,4),ncol=2)
D=matrix(c(5,7,6,8),ncol=2)
C*D
```

```
## [,1] [,2]
## [1,] 5 12
## [2,] 21 32
```

Warning: pay attention to dimensions of matrices

 Note: element-wise multiplication of two matrices also requries that dimensions (i.e. number of rows and columns) match:

```
E=matrix(c(1,3,2,4),ncol=1)
F=matrix(c(5,7,6,8),ncol=2)
E*F
```

Error in E * F: non-conformable arrays

- E is a 1 × 4 matrix.
- F is a 2×2 matrix.

-We want to multiply the element in the first row and first column of *E* with the element in the first row and the first column of *F* That's okay.

- ...
- We want to multiply the element in the third row and first column of E with the element in the third row and the first column of F. That's not doable, because F only has two rows.

Matrix multiplication

Consider again the following two matrices

$$C = \begin{bmatrix} 1, 2 \\ 3, 4 \end{bmatrix} \text{ and } D = \begin{bmatrix} 5, 6 \\ 7, 8 \end{bmatrix}$$

• Let's now consider the product of matrices *E* and *D*:

$$CD = \begin{bmatrix} 1 \times 5 + 2 \times 7, 1 \times 6 + 2 \times 8 \\ 3 \times 5 + 4 \times 7, 3 \times 6 + 4 \times 8 \end{bmatrix}$$

and in R:

```
C=matrix(c(1,3,2,4),ncol=2)
D=matrix(c(5,7,6,8),ncol=2)
C%*%D
```

```
## [,1] [,2]
## [1,] 19 22
## [2,] 43 50
```

Warning: pay attention to dimensions of matrices

 Note: matrix multiplication of two matrices requries that the number of rows in the left hand side matrix correspond to the number of columns in the right hand side matrix.

```
E=matrix(c(1,3,2,4),ncol=1)
F=matrix(c(5,7,6,8),ncol=2)
E%*%F
```

Error in E %*% F: non-conformable arguments

- E is a 1×4 matrix.
- F is a 2×2 matrix.

-We want to multiply the elements of the first row in matrix E to the elements of the first column of matrix F, but the former has one element and the latter has two elements!

Some special matrices

A 0-matrix (all entries are zero):

```
matrix(0, nrow = 2, ncol = 2)
## [,1] [,2]
```

```
## [1,] 0 0
## [2,] 0 0
```

• A J matrix (all entries are 1s):

```
matrix(1, nrow = 2, ncol = 2)
```

```
## [,1] [,2]
## [1,] 1 1
## [2,] 1 1
```

A matrix where all entries outside the diagonal are zero:

diag(c(1,2,3))

```
## [,1] [,2] [,3]
## [1,] 1 0 0
## [2,] 0 2 0
## [3,] 0 0 3
```

Some special matrices

diag(c(1,1))

• The identity matrix: a diagonal matrix where all elements in the diagonal are 1.

```
## [,1] [,2]
## [1,] 1 0
## [2,] 0 1
```

• An Identity matrix satisfies IA = AI = A, where A is a matrix.

```
A<-matrix(c(1,3,4,5),ncol=2)
I<-diag(c(1,1))
A%*%I
```

```
## [,1] [,2]
## [1,] 1 4
## [2,] 3 5
```

We can also apply diag() on matrix to extract the diaognal

```
A=matrix(c(1,3,4,5),ncol=2)
diag(A)
```

```
## [1] 1 5
```

The inverse of a matrix

- Let A be a n x n matrix (a square matrix, because the number of rows equals the number of columns).
- Let B be $n \times n$ matrix which multiplied by matrix A gives the identity matrix:

$$AB = BA = I$$

- The matrix B is called A's inverse, $B = A^{-1}$.
- Finding the inverse matrix is numerically complicated. But luckily we can ask R to do it for us by means of the solve() function:

```
A=matrix(c(1,3,4,5),ncol=2)
solve(A)
```

```
## [,1] [,2]
## [1,] -0.7142857 0.5714286
## [2,] 0.4285714 -0.1428571
```

let's test it:

solve(A)%*%A

```
## [,1] [,2]
## [1,] 1 -4.440892e-16
## [2,] 0 1.000000e+00
```

The determinant of a matrix

- A matrix is not invertible if the determinant is zero (the matrix is then called singular).
- The determinant of matrix A is written as det(A) or |A|.
- For a 2x2 matrix, the determinant is defined as

$$det(A) = |A| = \begin{vmatrix} a, b \\ c, d \end{vmatrix} = a \times d - b \times c.$$

- So for the A matrix defined earlier it is given by:

$$det(A) = \begin{vmatrix} 1,3\\4,5 \end{vmatrix} = 1 \times 5 - 3 \times 4 = -7.$$

A=matrix(c(1,3,4,5),ncol=2) det(A)

[1] -7

Combining matrices

Column bind: cbind(A,B,..)

• combines matrices A, B, .. horizontally (binds the columns).

```
A=matrix(c(1,3,4,5),ncol=2)
B=matrix(c(1,3,4,5),ncol=2)
cbind(A,B)
```

```
## [,1] [,2] [,3] [,4]
## [1,] 1 4 1 4
## [2,] 3 5 3 5
```

Row bind: rbind()

• combines matrices A, B, .. vertically (binds the columns).

rbind(A,B)

```
## [,1] [,2]
## [1,] 1 4
## [2,] 3 5
## [3,] 1 4
## [4,] 3 5
```

Application: ordinary least squares

 Let's return to our data and estimate the following model using Ordinary Least Squares (OLS):

$$test_year_6_i = \beta_0 + \beta_1 parental_lincome + \beta_2 summercamp_i + e_i$$

• We can achive this with lm() function in R:

```
mydataset<-read_csv("example_data1.csv")
my_lm<-lm(test_year_6~parental_lincome+summercamp,data=mydataset)
summary(my_lm)</pre>
```

- We specify the model to estimate on the form: y~x1+x2+.. (R automatically adds a constant).
 - We specify the data object to use using data=...
- We store the result of fitting the model using OLS in the object called my_lm
- We use the summary() function to summarize the results.

Application: ordinary least squares

Let's check the output

```
mydataset<-read_csv("example_data1.csv")
my_lm<-lm(test_year_6~parental_lincome+summercamp,data=mydataset)
summary(my_lm)</pre>
```

```
##
## Call:
## lm(formula = test_year_6 ~ parental_lincome + summercamp, data = mydataset)
##
## Residuals:
##
       Min
                 1Q Median
                                  30
                                         Max
## -0.95406 -0.28911 0.00042 0.26314 1.34439
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                 -7.82804 0.42003 -18.64 <2e-16 ***
## parental_lincome 0.66349 0.02821 23.52 <2e-16 ***
## summercamp 0.55130 0.03941 13.99 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.4022 on 488 degrees of freedom
## Multiple R-squared: 0.7059, Adjusted R-squared: 0.7047
## F-statistic: 585.6 on 2 and 488 DF, p-value: < 2.2e-16
```

Application: Ordinary Least Squares

Manual OLS using R

[1] "spec_tbl_df" "tbl_df"

- Let's try to manually reproduce these results from lm() using the matrix tools we iust covered.
- We know that the OLS estimator in matrix form is given by:

$$\hat{\beta} = (X^T X)^{-1} X^T y$$

 We know how to find the inverse, how to multiply matrices and how to transpose matrices in R. We are ready!

But

We first need to get the data from the dataset into the matrix.

```
mydataset <- read_csv("example_data1.csv")
class(mvdataset)
```

- "tbl" mydataset is a tibble (or a data frame), not a matrix.
- A matrix is homogeneous (just like vectors created with c()).
- Data frames allow for a mix of types (integer, double, character).

"data.frame"

From data frame to matrix: method 1

• We can access a specific column of a dataframe using the \$ symbol:

```
mydataset$test_year_6

## [1] 3.0968914 1.7556519 2.5691076 2.9608262 3.5352431 1.9390220 1.4565211 0.8530146 1.9422049 2.3242188 4.0378935 2.94128

## [60] 2.6405047 1.9769918 3.0799881 1.9301455 2.6791417 3.0750632 2.6380266 2.2997583 1.9301881 2.3110535 3.904595 2.73011

## [119] 2.1362477 1.9975301 1.9805466 3.2193071 1.7284275 1.3998921 1.8137896 2.5268532 2.4869136 1.8184657 2.8640394 3.46774

## [178] 2.2438199 2.7951478 2.3820597 3.3607016 1.9551251 2.2914024 2.5629094 2.5516064 2.4793032 1.7563591 2.6096906 2.19654

## [237] 2.8029133 2.6756473 3.0413707 1.9748095 3.8084274 1.5147583 3.2044558 2.7468996 2.1079639 1.9867444 2.7115403 2.12955

## [296] 3.2880623 2.9644639 2.1247169 2.8076262 2.2058520 1.3455721 1.8300033 2.7072177 2.7233555 1.2591008 2.3801160 3.11343
```

[343] 2.9544985 1.9224016 2.5383838 2.4922356 3.1961359 3.2343892 4.0994714 2.9938602 2.8084186 3.0216950 2.6169923 3.24302
[414] 2.2764986 1.7132658 2.3578819 2.5405297 1.6079901 1.4885995 2.6985074 2.2916481 2.6920764 1.5655158 2.4127587 2.35459
[473] 3.6784746 2.412565 2.4748137 4.0057407 1.1091630 3.5154961 2.4298711 2.2944919 3.4194154 2.1559040 2.2093878 3.13277

let's save data in a column vector called y:

mydataset<-read_csv("example_data1.csv")</pre>

```
mydataset<-read_csv("example_data1.csv")
y<-matrix(mydataset$test_year_6,ncol=1)</pre>
```

 Note that we also could use the [] subsetting method, but this requires us to know the order of the columns in the dataset.

From data frame to matrix: method 2

• We can convert a data frame to a matrix using the as.matrix() function:

[1] "matrix" "array"

let's check the content of X by printing the first five rows and all columns:

```
print(X[1:5,])
```

```
## parental_lincome summercamp constant
## [1,] 15.26362 1 1
## [2,] 13.96494 0 1 1
## [3,] 15.06514 1 1
## [4,] 15.32181 1 1
## [5,] 15.71261 1 1 1
```

Let's find the OLS estimates

• The point-estimates of the OLS estimator are given by:

$$\hat{\beta} = (X^T X)^{-1} X^T y$$

Let's translate this into R.

```
# load dataset
mydataset<-read_csv("example_data1.csv")
# extract the dependent variable and store as column vector y
y<-matrix(mydataset$test_year_6,ncol=1)
# extract the covariates to include and create a constant and as matrix X
X<-mydataset\%'\%
    select(parental_lincome, summercamp) %\%
    mutate(constant=1) %\%
    as.matrix()
# implement OLS formula
betahat<-solve(t(X)%*\%X)%*\%t(X)%*\%y
# show betahat vector
print(betahat)</pre>
```

[,1]
parental_lincome 0.6634867
summercamp 0.5513035
constant -7.8280434

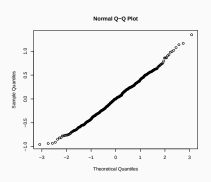
Estimated coefficients are identical to the results obtained with lm()!

Application: Ordinary Least Squares

Fitted values

- the fitted values are given by $X\hat{\beta}$.
- the residuals are given by $y X\hat{\beta}$.
- we can create a q-q plot of the residuals using base graphics:

```
#calculate residuals
residuals<-(y-X%*% betahat)
# create a q-q plot
qqnorm(residuals)</pre>
```



Matrices in R - summary

- Create column vector A: A<-c().
- Create matrix A: A <- matrix().
- Convert B to a matrix and store it in A: A<-as.matrix(B).
- Transpose matrix A: t(A).
- Inverse of matrix A: solve().
- Element-wise multiplication of matrix A and B: A*B.
- Matrix multiplication of matrix A and B: A%*%B.
- Extract the column named col1 from data frame df: df\$col1.
- OLS estimator: betahat=solve(t(X)%*%X)%*%t(X)%*%y.

4. Functions, control structures and loops in $\ensuremath{\mathsf{R}}$

Defining functions

What is a function?

- a set of R statements that perform a task given a set of provided arguments.
- Example: lm(test_year_6~parental_lincome+summercamp,data=mydataset)
 - The function lm() estimates coefficients of a linear model.
 - The model is provided as the first argument (test_year_6~parental_lincome+summercamp)
 - The dataset is provided as a second (named) argument (mydataset).

User defined functions

- We can easily create our own functions in R.
- The syntax is as follows:

```
function_name <- function(arg_1, arg_2, ...) {
   Function body
}</pre>
```

The function consists of four parts:

- 1. The function name.
- 2. The arguments (placeholders for settings, datasets etc).
- 3. The function body (a collection of statements to carry out using the arguments provided).
- 4. Return value (the last expression of the function)

Our first function

Let's define a function

- 1. name: Hansfunction
- 2. arguments: x and y
- 3. function body: z=x*y
- 4. return: return(z)
- in R:

```
Hansfunction <- function(x, y) {
   z<-x*y
   z
}</pre>
```

Let's try the function

Hansfunction(3,6)

[1] 18

• What if we forget to state an argument?

Hansfunction(3)

 $\mbox{\tt \#\#}$ Error in Hansfunction(3): argument "y" is missing, with no default

Default values

Let's define a function with default values

- To avoid such cases, we can specify **default** values:
- When defining the function, we set the arguments equal to their default values.
- Let's define that function.

```
Hansfunction <- function(x=2, y=2) {
   z<-x*y
   z
}
Hansfunction(3)</pre>
```

```
## [1] 6
```

 This works well, but if you accidentially forgot an argument? It would be nice with a warning.

Control structures

Control structures

- We can use **control structures** to control the statements executed by our function.
- Here is an example of a control structure in plain English:

```
if logical test evaluates to true do the following statements to execute if TRUE
else do the following statements to execute if not TRUE
```

• and in terms of R syntax:

```
if (logical test){
    }
    else{
}
```

- A logical test is a statement that evaluates to TRUE or FALSE, for example:
 - "5 is greater than 3" The statement is TRUE
 - "3 is greater than 5" The statement is FALSE
- Control statements can also be used outside functions (in scripts, loops etc).

Control structures in Hansfunction()

```
Hansfunction <- function(x=2, y=3) {
  if (missing(x)|missing(y)){
    print("Warning: Not all arguments provided. Default values used.")
  }
  else{
    print("Well done, you specified all arguments!")
  }
    z<-x*y
    z
}
Hansfunction(3)</pre>
```

- ## [1] "Warning: Not all arguments provided. Default values used."
- ## [1] 9
 - here we use the function missing() to test whether the argument missing.
 - the | corresponds to "or" (the logical expression evaluates to true if x OR y are missing).
 - We can use the symbol "&" if we only want the expression to evaluate to true if both are x and y are missing.

Local vs global

- Global R objects are accessible from anywhere.
- Local R objects only exist in local environment.

```
assign_to_z <- function(x) {
    z<-x
}
z=5
assign_to_z(3)
print(z)</pre>
```

[1] 5

- What is going on? Inside a function is a local environment.
- Changing and creating objects within local environments does affect global environments.
- Unless we explicitly tell R to do so:

```
assign_to_z <- function(x) {
    z<<-x
}
z=5
assign_to_z(3)
print(z)</pre>
```

[1] 3

Note the use of <<- (the super assignment operator).

Our own Im() function

Let's build our own Im() function

 We now use the tools from the matrix section and combine them with the function definitions to create our own Im function.

```
mylm <- function(y,x,data) {
  # specify dataset
  df<-data
  # extract the dependent variable and the covariates
  yvar<-df%>%select(y)%>%as.matrix()
  Xvar<-df%>%select(x)%>%mutate(constant=1)%>%as.matrix()
  # implement OLS formula and return betahat vector
  solve(t(Xvar)%*%Xvar)%*%t(Xvar)%*%yvar
# load data
mydataset<-read csv("example data1.csv")</pre>
# tru our new function
mylm(x=c("parental_lincome", "summercamp"), y="test_year_4", data=mydataset)
## Note: Using an external vector in selections is ambiguous.
## i Use 'all_of(y)' instead of 'y' to silence this message.
## i See <https://tidyselect.r-lib.org/reference/faq-external-vector.html>.
## This message is displayed once per session.
## Note: Using an external vector in selections is ambiguous.
## i Use 'all_of(x)' instead of 'x' to silence this message.
## i See <a href="https://tidyselect.r-lib.org/reference/faq-external-vector.html">https://tidyselect.r-lib.org/reference/faq-external-vector.html</a>.
## This message is displayed once per session.
##
                  test_year_4
## parental_lincome 0.6249679
```

For loop

- A loop repeats a set of statements.
- The number of times the statements are repeated is stated in the loop header.
- The set statements are provided in the loop body.
- Example:

```
for (x in 1:3){
  print(x)
}
```

- ## [1] 1
- ## [1] 2
- ## [1] 3
 - The loop header for (x in 1:3){ states that the loop should be repeated 3 times:
 - 1. Once where x has the value 1
 - 2. Once where x has the value 2
 - 3. Once where x has the value 3.
 - The loop header states that in each loop iteration the statment print(x) should be executed.

While loop

- We can also create a loop that repats itself until a certain condition is violated.
- This is called a while loop.
- The loop body statements are repeated until the while condition is violated.
- Example:

```
x=1
while (x<5){
  print(x)
  x=x+1
}</pre>
```

- ## [1] 1
- ## [1] 2
- ## [1] 3
- ## [1] 4
 - the object x is initiated with a value of 1.
 - the loop is repeated until x<5 evaluates to false.
 - in every iteration we:
 - print the value of x.
 - add the value of 1 to x.

Maximum likelihood: find the parameters that maximize the likelihood that we observe what we've observed, given an assumed functional form and distribution.

- Application: estimate the probability that a child participates in the summer school using a probit model.
- Benchmark: R's built-in function

```
##
## Call:
## glm(formula = summercamp ~ parental_lincome, family = binomial(link = "probit"),
##
      data = mvdataset)
##
## Deviance Residuals:
      Min
               1Q Median
                                        Max
## -2.2385 -1.0561 0.5512 1.0291 1.8498
##
## Coefficients:
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept) -11.93926 1.44268 -8.276 <2e-16 ***
## parental_lincome 0.79602 0.09538 8.346 <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
```

The likelihood for a single observation:

$$L(\beta; y_i, x_i) = [\Phi(x_i'\beta)]^{y_i} [1 - \Phi(x_i'\beta)]^{1 - y_i}$$

 Observations are assumed to be iid, we can therefore write the likelihood of the entire sample as the product of the individual likelihoods:

$$L(\beta; y, X) = \prod_{i=1}^{N} [\Phi(x_t'\beta)]^{y_i} [1 - \Phi(x_t'\beta)]^{1-y_i}$$

The log-likelihood is then given by:

$$I(\beta; y, X) = \sum_{t} \left(y_t \ln \Phi(x_t'\beta) + (1 - y_t) \ln \left(1 - \Phi(x_t'\beta) \right) \right)$$

- R implementation
 - Φ() is the cumulative distribution function of the standard normal distribution, which we implement in R with pnorm().
 - sum() computs the sum.
 - We can therefore implement the above in R as: l<-sum(y*log(pnorm(xb))+(1-y)*log(1-pnorm(xb)))</p>

• The log-likelihood for $\beta = [-11.9, 0.79]$ (the values R found for us).

```
# load data
df<-read_csv("example_data1.csv")
# y variable
y<-df%>%select(summercamp)%>%as.matrix()
# x variable
X<-df%>%select(parental_lincome)%>%mutate(constant=1)%>%as.matrix()
# xb (note constant is given last)
xb<-X%+%c(0.79,-11.9)
# log likelihood
l<-sum(y*log(pnorm(xb))+(1-y)*log(1-pnorm(xb)))
# return value
1</pre>
```

- ## [1] -300.0516
 - Okay, but how do we know this is maximized? Let's evaluate the log likelihood value for various values of beta. To do this we:
 - 1. Wrap the likelihood expression in a function.
 - 2. Loop over the function and use different values.

1. Our likelihood function

```
my_loglikelihood <- function(y,x,beta,data) {
  # specify dataset
  df<-data
  # extract the dependent variable and the covariates
 vvar<-df%>%select(v)%>%as.matrix()
  Xvar<-df%>%select(x)%>%mutate(constant=1)%>%as.matrix()
  # heta
  betavec=beta
  # xb (note constant is given last)
 xb<-Xvar%*%betavec
  # log likelihood
 1<-sum(yvar*log(pnorm(xb))+(1-yvar)*log(1-pnorm(xb)))</pre>
  # return (explicitly tell R to return this object)!
 return(1)
# load data
mydataset<-read_csv("example_data1.csv")</pre>
# try our new function
my loglikelihood(x="parental lincome", y="summercamp",
                 beta=c(0.79,-11.9) ,data=mydataset)
```

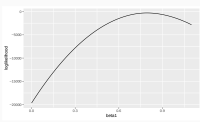
[1] -300.0516

2. Loop over values

• Simplification: keep β_0 constant at -11.9 and only change β_1 :

```
# loop over values of beta1
for (i in seq(0,1,by=0.2)){
  1<-(my loglikelihood(x="parental lincome",y="summercamp",</pre>
                 beta=c(i,-11.9),data=mvdataset))
  print(paste("For beta1=" ,i, " the log-likelihood is: ",1,".", sep=""))
  [1] "For beta1=0 the log-likelihood is: -19664.9657835808."
   [1] "For beta1=0.2 the log-likelihood is: -11137.5254933262."
  [1] "For beta1=0.4 the log-likelihood is: -5093.99325748709."
   [1] "For beta1=0.6 the log-likelihood is: -1493.09962685655."
## [1] "For beta1=0.8 the log-likelihood is: -301.062735618683."
## [1] "For beta1=1 the log-likelihood is: -1503.16885413735."
 Okav. that works, but 5 values are a bit boring. Let's try more!
```

• Let's loop over more values, store all results in a data frame and show the likelihood in a chart as a function of β_1 :



- The chart indicates that the maximum value of the log-likelihood function could be around 0.79 (as the R built-in function suggest).
- But how do we find the exact values? And how about β_0 ?
- We use a built-in optimizer.
 - A function to maximize or minimize to mimize an expression.
- One such function is optim(), In optim() we specify:
 - par: starting values for the parameters.
 - fn: the function to maximize (or minimize).
 - control a control parameter (fnscale=-1 is a scaling parameter that we apply on the values. By setting it to -1 it becomes a maximization problem).
 - ...: options passed on to the function in fn.

```
## $par
## [1] 0.7960093 -11.9392406
## $value
## [1] -299.6738
## $counts
## function gradient
## 89 NA
##
## $convergence
## [1] 0
##
## $message
```

Functions, control structures, loops - summary

• We define functions using the following syntax:

```
function_name <- function(arg_1, arg_2, ...) {
  function body
}</pre>
```

• We control the flow of our function using control structure:

```
if (logical test){
   action to do if logical test evaluates to true
}
else {
   action to do if logical test evaluates to false
}
```

We repeat statements using loops

```
for (x in range){
  action to repeat for all values in range
}
```

We can combine these tools and implement a maximum likelihood estimator.



Please send suggestions for improvements and notes about identified mistakes to h.h.sievertsen@bristol.ac.uk.

Thanks.