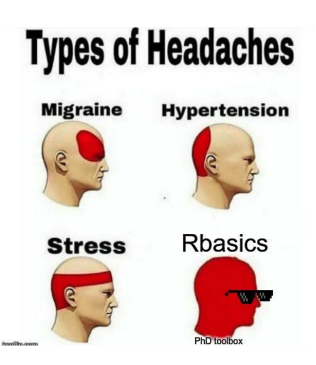
# Rbasics PhD toolbox

Matteo Chialva Martino Adamo

## Course organization

```
RBASICs
Day 1
      Part I - Operative Introduction to R
      Part II - Spread Sheets
      Part III - Manipulating data with dplyr (may shift in part to day2)
Day 2
      Part IV - Text strings
      Part V - Tips and Tricks
      Part VI - Base graphics in R
```

## Part I Operative introduction to R



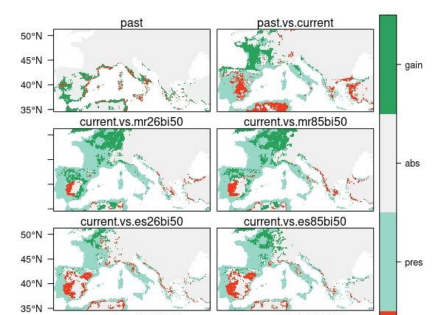
#### Part 1 contents

- why R instead of other software (excel, SPSS..)
- R is not only a statistics software
- using R through RStudio: the main features
- installing and updating R and RStudio software
- base and advanced packages installation using external repositories (BioConductor, github..).
- systems dependencies and packages
- R objects type: vectors, matrices, dataframes, lists & functions
- data import and export (read.table; write.table; load; WriteXLS)
- practical tips for organizing scripts, files and folders
- "don't panic!": guidelines to face issues during a data analysis session in R (error messages and warnings, manuals, tutorials, forums, reference sites)

#### teachers for a week

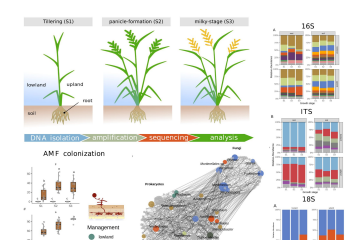
#### Dr. Martino ADAMO

Postdoc researcher since 2018, I work on plants <u>diversity</u> <u>conservation</u> and <u>orchid mycorrhizas</u> using bio-molecular tools and boring statistical model to study traits influence on species <u>spatial distribution</u>.



#### Dr. Matteo CHIALVA

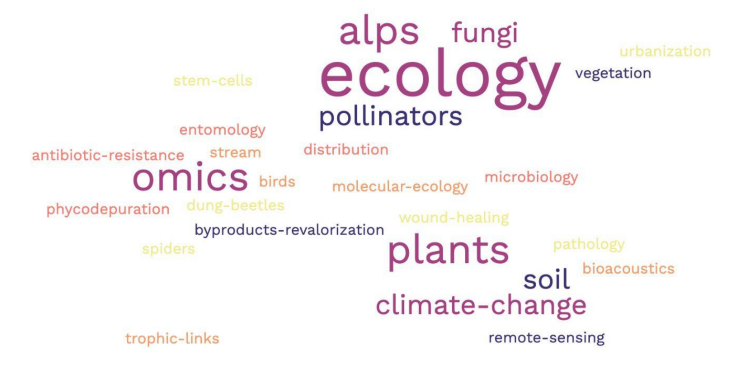
He is post-doctoral resercher since 2017 and his research focuses on <u>plant-microbes interactions</u> in model crop species by using multi-omics approaches from transcriptomics to metagenomics. By coupling these tools with systems biology and biostatistics he is interested in linking <u>soil microbiota diversity</u> and functioning to plant responses and ecosystem services.



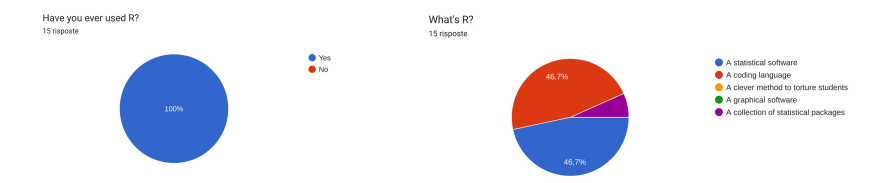
#### students forever

PhD is the first step of academic career: if you want to continue you are going to study every single day of your working life.

We asked you to explain your study field in three keywords and this is results:



## What's R?

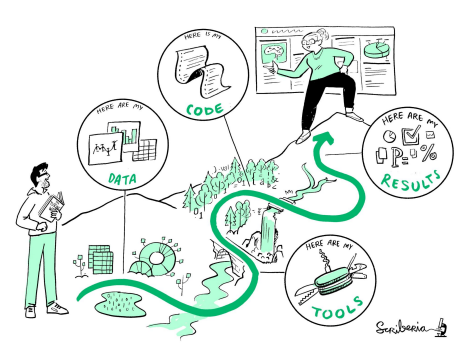


All of you know that R exist and have already used a time in their life

but....less then half of you know what R really is!

R per se is a programming environment for statistics and graphics, not a simple standalone software. Much more similar to a coding language rather than a stat. software

#### why R instead of other software



1. R is free and open-source

2. R can do almost everything. No seriously, everything.

3. R can address many of the challenges of performing reproducible research

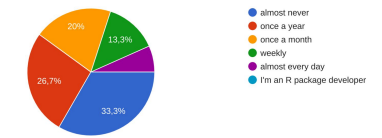
4. Learning R will make you a more attractive candidate

#### What's R?

Most of you rarely used R

How frequently do you use R?

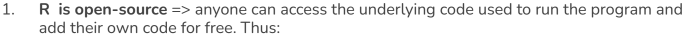
15 risposte



#### Why you should use it **every day**:

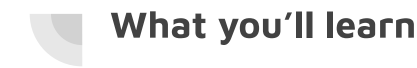
- open source and cross platform
- active community of developers: i.e. access to the state-of-the-art methods
- extensive documentation everywhere (books, online, forums etc..)
- great graphics capabilities: you can manage all the type of plots you can imagine (and mainly those that you could not!)

#### why R instead of other software



- a. will always be able to perform the newest statistical analyses as soon as anyone thinks of them;
- will fix its bugs quickly and transparently; and
- c. has brought together a community of programming and stats nerds (a.k.a., useRs) that you can turn to for help.
- 2. **R can do almost everything**. No seriously, everything.
  - a. basic stats
  - b. math
  - c. frequentist and Bayesian statistics.
  - d. excel suffs
  - e. it saves money!
  - f. LaTeX and Word stuffs
  - g. Multicore processing
  - h. Phylogeny
  - i. genomics
  - i. make fun
  - k. ..
- 3. R can address many of the challenges of performing **reproducible research**
- 4. A final reason you should become a useR is that R is increasingly being used as an industry standard in the realm of data analytics, also known as "data science." Learning R will make you a more attractive candidate if you apply for non-academic jobs and academic positions.



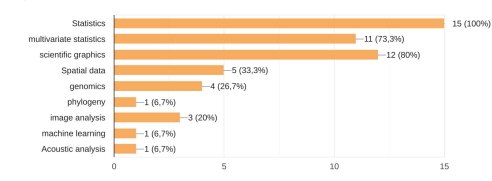


With R you can do analyses in all the scientific disciplines.

Here you'll learn R and its main functions to deal with tables (any type of data) and graphics.

You'll not learn statistics!

Which kind of data analysis do you usually implement or you plan to deal with? 15 risposte



Our aim is to provide you universal tools to deal with your data in a reproducible way



#### Reproducible research

#### What is reproducibility?

The possibility to reproduce someone else results. It's part of the scientific method (it should be...)

How can you ensure reproducibility?

One of the best way is to use **CODE** since it is universal and can be easily shared and documented

Reduce (and possibly avoid) manual data manipulation steps (e.g. copy-paste).

- prone to introduce errors
- time inefficient
- impossible to reproduce

If you cannot avoid that you have to document what has been done and what was the purpose.

#### installing R and RStudio

Almost everyone run R through RStudio GUI (graphic user interface) which is a software which helps you to write in R.

- 1. Install R <a href="https://cran.r-project.org/">https://cran.r-project.org/</a>
- 2. Install RStudio (free version) <a href="https://posit.co/downloads/">https://posit.co/downloads/</a>

Note: in both cases installation procedure dependecies on your operating system.

(a dependency is additional code that a programmer wants to call)

## updating R and RStudio

Periodically you would have to update both R and/or RStudio.

1. Update R - different options available

The most effective way is to completely remove the previous R version and install the new one.

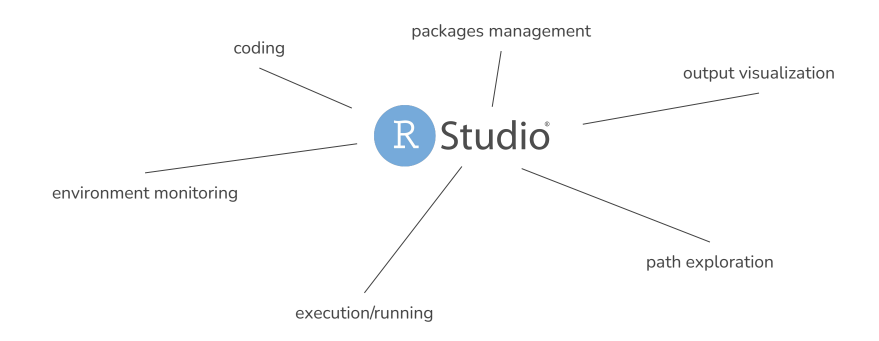
Note: you have to fully re-install all the packages or move the old packages in the new directory.

2. Update RStudio

Download and install the new version (removal of the previous version is highly suggested!)

#### using R through RStudio: the main features

RStudio is an integrated development environment for R



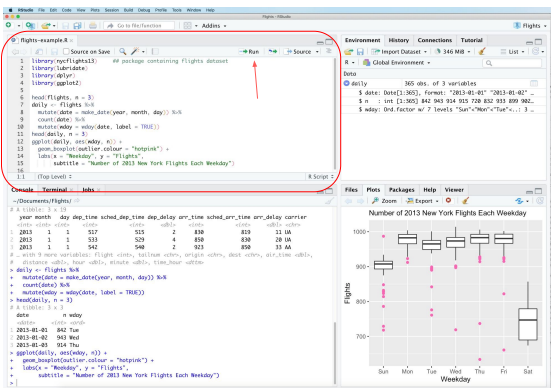
#### using R through RStudio: the main feature

#### coding

here you can write your code

90% of the time on Rstudio you will work here or here:





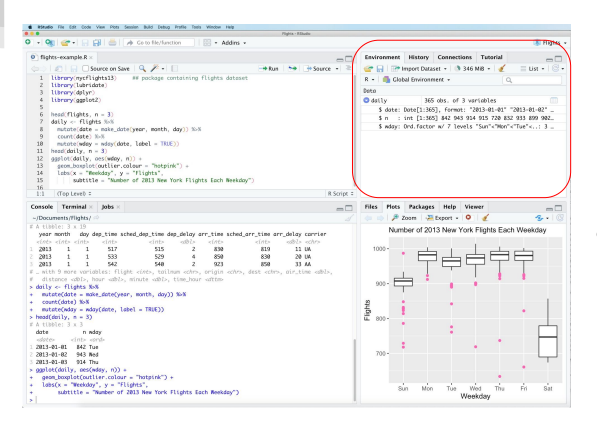


RStudio File Edit Code View Plots Session Build Debug Profile Tools Window Help O - On On File/function Flights @ flights-example.R Environment History Connections Tutorial ⟨□□⟩ | Ø□ | □ Source on Save | Q 
Ø → | □ Run Source - = Import Dataset - 346 MiB - List → | C -1 library(nycflights13) ## package containing flights dataset R - Global Environment -2 library(lubridate) Data 3 library(dplyr) 4 library(ggplot2) o daily 365 obs. of 3 variables \$ date: Date[1:365], format: "2013-01-01" "2013-01-02" ... head(flights, n = 3)\$ n : int [1:365] 842 943 914 915 720 832 933 899 902... daily <- flights %>% \$ wday: Ord.factor w/ 7 levels "Sun"<"Mon"<"Tue"<...: 3 ... mutate(date = make date(vear, month, day)) %>% mutate(wday = wday(date, label = TRUE)) 11 head(daily, n = 3) ggplot(daily, aes(wday, n)) + geom\_boxplot(outlier.colour = "hotpink") + labs(x = "Weekday", y = "Flights", 15 subtitle = "Number of 2013 New York Flights Each Weekday") (Top Level) \$ R Script \$ 1:1 Console Terminal × Jobs Files Plots Packages Help Viewer -0 ~/Documents/Flights/ Zoom 
 Zexport 
 Number of 2013 New York Flights Each Weekday year month day dep\_time sched\_dep\_time dep\_delay arr\_time sched\_arr\_time arr\_delay carrier <int> <dh1> 819 11 UA 529 20 UA # \_ with 9 more variables: flight <int>, tailnum <chr>, origin <chr>, dest <chr>, air\_time <dbl>, # distance <dbl>, hour <dbl>, minute <dbl>, time\_hour <dttm> > daily <- flights %>% + mutate(date = make\_date(year, month, day)) %>% Flights count(date) %>% + mutate(wday = wday(date, label = TRUE)) > head(daily, n = 3) # A tibble: 3 x 3 700 -2013-01-02 943 Wed 2013-01-03 914 Thu > ggplot(daily, aes(wday, n)) + geom\_boxplot(outlier.colour = "hotpink") + labs(x = "Weekday", y = "Flights", subtitle = "Number of 2013 New York Flights Each Weekday") Weekday

**console** this is actually R

hybrid function coding:: ::running:: ::results viz

#### using R through RStudio: the main feature



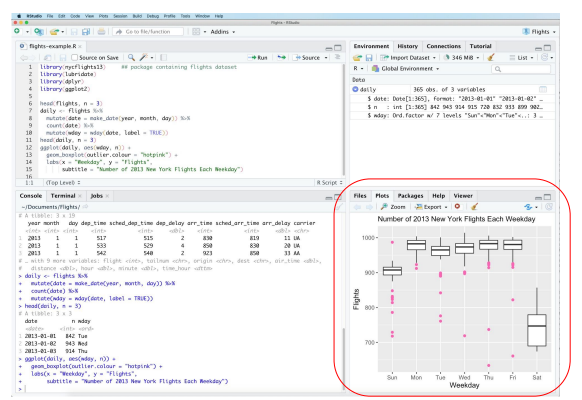
#### environment

this it the best Rstudio intuition

R commands work on objects

here you can see the objects you're working on

#### using R through RStudio: the main feature



#### plots and explorer

here you see graphical outputs, help pages and your packages

moreover you can explore your folders and files

## working directory

wd is where you're working and where R looks for files to import or to save outputs ... default wd is into the R folder in your home, we suggest to set as wd the folder where your input files are stored.

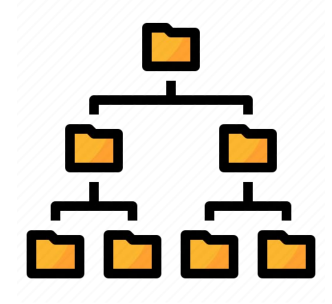
setwd("your path")

getwd()

Example path formats in different OS.

/ubuntu/unix/macos/your\_dir/your\_filename

"C:\Users\your\_folder\your\_filename"







• **CRAN**: it is a network of ftp and web servers around the world that store identical, up-to-date, versions of code and documentation for R.

```
>install.package("pkg.name") OR via RStudio interface
```

 Bioconductor: develop, support, and disseminate free open source software that facilitates rigorous and reproducible bioinfo analysis

```
>BiocManager::install(c("package1", "package2"))
```



• From external repositories, e.g. from GitHub: it is an online software development platform. It's used for storing, tracking, and collaborating on software projects



- using **devtools**: it makes package development easier by providing R functions that simplify and expedite common tasks

```
>devtools::install_github("repository_name")
```

- install CRAN and special packages we'll use most of these packages later on..
  - install a package from CRAN:
    - 'devtools'
    - 'BiocManager'
    - 'stringr'
    - 'reshape'
    - 'tidyverse'

>install.packages(c("devtools", "BiocManager", ...))

- install a package from GitHub (https://github.com/jfq3/QsRutils)
  - https://github.com/jfq3/QsRutils

Github packages are installed through the devtools::install\_github() function

> install\_github("jfq3/QsRutils")

- install a package from Bioconductor (Biostrings)
  - https://www.bioconductor.org/packages/release/bioc/html/Biost rings.html

Bioconductor packages are installed through the BiocManager::install() function

>BiocManager::install("Biostrings")

If for some reason installation fails R reports you some errors Let's check:

- they are in the list of the installed packages
- they can be loaded. Use the function library()

## Few notes on programming

- **R** is the most simple coding language available
- You don't need to be an experience programmer to use R language
- as every language, R has its own grammar
- Let RStudio help you in writing in R (as Word helps you writing in another language)
- Don't panic, but...you'll need to memorize a lot of functions
- most of R functions are simple and intuitive, the best way is to practice!

## Few notes on programming

- coding is case sensitive (e.g. **object** and **Object** are not the same thing!)
- "#" symbol in general excludes lines that the language runs. It is used to comment-out lines with alternative/inactive code or to annotate your script.
- text strings are in quotes: e.g. "Object" is the string text name of Object
- objects names cannot contain spaces or mathematical operators
- possibly keep objects and functions name different!
- objects are created, replaced, filled with the symbol "<-"</li>
   object <- "I'm an object"</li>
- "<-" and "=" are alternatives

#### data type

• numeric (num) = numbers

```
num [1:91] 1 1.1 1.2 1.3 1.4 1.5 1.6 1.7 1.8 1.9 ..
```

• factor (factor) = categories, ordinal numbers, boolean

```
Factor w/ 2 levels "due", "uno": 2 1
```

• character (char) = text

```
chr [1:8] "A" "A" "B" "B" "A" "A" "B" "B"
```

• integer (int) = integer numbers

```
int [1:10, 1:5] 41 18 25 12 13 23 42 10 21 2 ...
```

• logical (logi) = TRUE/FALSE values (boolean data type)

```
logi [1:5] TRUE FALSE FALSE FALSE TRUE
```

Boolean are all two-step variables: True-False Yes-No 1-0

## object class

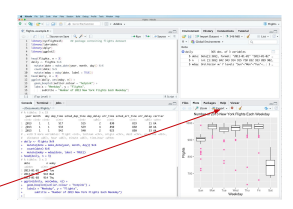
atomic vectors, matrices, dataframes, lists & functions + special features

#### R object class: functions

> install.packages() is a R function.

R function are **ALWAYS** followed by brackets

R functions are grouped in **packages**.



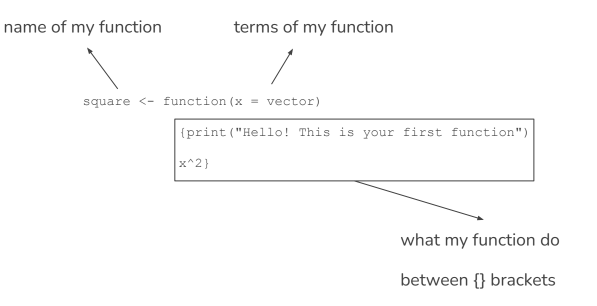
each function is more or less explained in a **help page**:

#### <u>Usaqe</u>



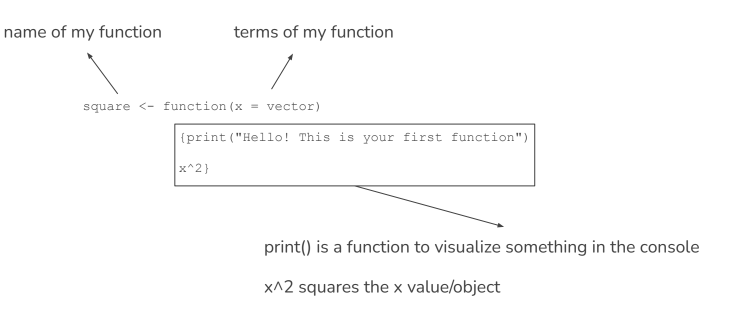
### R objects class: functions

Everybody in R can write functions. Just a naive example to understand their grammar:



### R objects class: <u>functions</u>

Everybody in R can write functions. Just a naive example to understand their grammar:

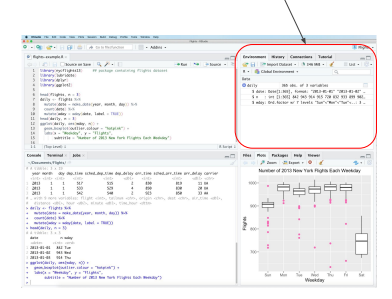


#### R objects class: functions

After the previous step you will have new object (a function in this case) in the Rstudio object list

so you can use it!

```
> square(12.567)
[1] "Hello! This is your first function"
[1] 157.9295
```





#### R objects class: <u>functions</u>

Most of the time you don't need to write your own functions but to look for needed functions among available R packages.

#### Some shortcuts:

?factor # displays the function help page

args(factor) # prints the arguments the function can take

## objects class: vector

A sequence of elements (number, characters) assigned using the c() function

Vector can contain only one data type, otherwise R tries to convert the content to avoid loosing information.

#### simple vector

> my\_vector <- c("first\_element", "second\_element", number, ...)

#### named vector

> my\_vector <- c("name\_1"="first\_element", "name\_2"="second\_element", "name\_3"="second\_element", ...)

#### display vector elements name

> names(my\_vector)

Exercise: write a numeric vector in a R object named "my\_vector"

## objects class: vector

```
# Write a simple vector:
> colors<-c("red", "blue", "yellow", "orange", "green", "brown")
# Check vector length with length() function
> length(colors)
[1] 6
```

## objects class: vector

```
# type of data in vector
> class(colors)
[1] "character"
# str() gives an overview of the structure of an object and its elements
> str(colors)
chr [1:6] "red" "blue" "yellow" "orange" "green" "brown"
```

```
> unique(colors)
[1] "red" "blue" "yellow" "orange" "green" "brown"
# add elements to a vector
> c(colors, "purple") # add to the end of the vector
[1] "red" "blue" "yellow" "orange" "green" "brown" "purple"
> c("white", colors) # add to the beginning of the vector
[1] "white" "red" "blue" "yellow" "orange" "green" "brown"
```

some shortcuts to work with vectors

```
# sequence of numbers from x to y
> seq(1, 10, by=1)
> vector <- rep(c("A", "B"), 2, each=2) # repeat elements in vectors
[1] "A" "A" "B" "B" "A" "A" "B" "B"
> vector 1 <- sample(seq(1,30), 5)
                                                   # generate vector with sampled random
elements
[1] 22 8 7 18 28
```

### **Exercise:**

### Produce:

- sequence of number (from 3475 3500 at 0.5 steps)
- repeated sequence of factors (repeat the words cat, tree and dog two times each and generate a vector of length 18
- vector of two random numbers between 0 and 1 (considering two numbers after the decimal point)

### **Exercise:**

- sequence of number (from 3500 to 3475 at 0.5 steps)

```
> seq(3475, 3500, by=0.5)
```

- repeated sequence of factors (repeat the words cat, tree and dog two times each and generate a vector of length 18

```
> rep(c("cat", "tree", "dog"), 3, each=2)
```

- vector of two random numbers between 0 and 1 (considering two numbers after the decimal point)

```
>  sample (seq (0,1, by=0.01), 2)
```

## subsetting vectors

To extract values from vectors positional indices should be provided in square brackets []

```
> animals<-c("cat", "dog", "rabbit", "duck", "monkey", "fish")
> animals[2]
[1] "dog"
```

## subsetting vectors

# multiple elements can be selected

```
> animals[c(3,2)]
[1] "rabbit" "dog"

# conditional subsetting:
> numbers<-c(24, 5, 47, 6, 98, 2, 10, 144)
> numbers[numbers>60]
[1] 98 144
```

## operations with vectors

Sum two or more vectors:

```
> c(1,1,1,1)+c(1,1,1,1)
[1] 2 2 2 2
```

if vectors are not the same length, the shorter will be recycled

```
> c(1,2)+c(1,1,1,1)
[1] 2 3 2 3
```

Arithmetic operations on vectors applies element-wise

```
> c(1,1,1,1)*5
[1] 5 5 5 5
```

### **Exercise:**

- write a vector in an object called "num" made of 10000 numbers comprised in the interval between 2 and 20000
- count how many values >3000
- **sum** elements number 385, 1001 and 7521

#### **Solution**

```
# generate num vector
> num < -seq(2, 20000, by=2)
# check vector length
> length(num)
[1] 10000
# subset and count elements
> length(num[num>3000])
> [1] 8500
# sum elements number 385, 1001 and 7521
> num[385]+num[1001]+num[7521]
[1] 17814
> sum(num[c(385,1001,7521)])
[1] 17814
```

### object class: matrices

```
> matrix(data = sample(seq(1,50)), nrow = 10, ncol = 5, byrow = F)
     [,1] [,2] [,3] [,4] [,5]
[1,]
       2
         21
               37
                   30
                      43
[2,]
      12
          42 9
                       46
[3,]
       7 19
              29 26
                       40
[4,]
      17
          32 39 23
                       25
[5,]
      10
         27
               50 33 15
[6,]
       5 13 1 44
[7,]
      47
          24 6 22 34
[8,]
      28
         18 14
                  45 36
         48 16
[9,]
      11
                       49
[10,]
          31
             38
                   20
                      35
      41
```

Matrices are grouped NUMERIC vectors

# object class: matrices

Matrices are grouped NUMERIC vectors, thus you can create matrices starting from several different vectors.

#### Exercise:

- 1. create 3 different vectors of length 45
  - vector 1 should be created using seq
  - vector 2 should be created using rep
  - vector 3 should be created using sample
- # repeat 45, 45 times
  - # 45 random numbers between 1 and 100
- 2. build your own matrix named "matrix" merging the vectors with the rbind() function

## object class: matrices

- 1. create 3 different vectors of length 45
  - vector 1 should be created using seq

```
> vector 1<-seq(1,45)
```

- vector 2 should be created using rep # repeat 45, 45 times
  - > vector\_2<-rep(45, 45)
- vector 3 should be created using sample # 45 random numbers between 1 and 100
  - > vector 3<-sample(seq(1,100), 4\$)</pre>
- 2. build your own matrix named "matrix" merging the vectors with the **rbind()** function

```
> my_matrix<-rbind(vector_1, vector_2, vector_3)</pre>
```

## object class: dataframe

a dataframe is a <u>group of any type of vectors</u> (character, number, integer, factor)...dataframes are the most common object in R

Manually write a dataframe using data.frame() function

```
"colname_2"=sample(seq(1,30), 8),  # vector 2
          "colname 3"=seq(8,11.5, by=0.5))  # vector 3
> df
 colname 1 colname 2 colname 3
                 8.0
                 8.5
3
                9.0
      В 13
           30
                 9.5
5
       18
                10.0
           29
                10.5
           21
                11.0
8
      В
           10
                11.5
```

## object class: list

a list is a group of any type of objects (vector, matrix, data.frame)...function is list(), each object in the list can be identified by a name

```
> my list<-list("vector"=vector, "matrix"=matrix, "dataframe"=df)</pre>
> my list
$vector
[1] "A" "A" "B" "B" "A" "A" "B" "B"
$matrix
    ΑВ
[1,] 1 2
[2,] 1 2
[3,] 1 2
$dataframe
  colname 1 colname 2 colname 3
                           8.0
                        8.5
         Α
                  20
                      9.0
                  21
                        9.5
                   9
                         10.0
         Α
         Α
                  28
                         10.5
                          11.0
                  12
                          11.5
```

# object class

each object can be duplicated (or overwritten!)

```
df2 <- df
my_list2 <- my_list
each object can be removed by rm()
rm(my_list2)</pre>
```

### object class

```
# Find the type of object
> class(df)
[1] "data.frame"
# inspect object (find the data structure of a R object)
> str(df)
'data.frame': 8 obs. of 3 variables:
 $ colname 1: chr "A" "A" "B" "B" ...
 $ colname 2: int 13 22 18 23 10 29 8 7
 $ colname 3: num 8 8.5 9 9.5 10 10.5 11 11.5
```

# object class

```
# Inspect dimensions
> dim(df)  # applies to data.frame and matrices
[1] 8 3
> length(vector) # applies to vectors only
[1] 8
```

# object type: special objects

We are not here to teach specific packages, but keep in mind that several specialist package can use the "special objects" they are often lists built with specific parameters!

This is the case of graphics objects (ggplot...)

APE

**POPPR** 

PHYLOSEQ

DESeq

... and many other...

### row- and column names

Each columns or rows in a given object (vector, matrix, dataframe) can have a name.

```
# display column names: colnames()
```

```
> colnames(object)
```

# display row names: rownames()

```
> rownames(object)
```

The same function can be used to change or set row/column names:

```
> colnames(object)<-c("colname1", "colname2")</pre>
```

Columns and rows can be flipped by transposition using the function t()

```
> matrix
    [,1] [,2]
[1,]
[2,] 1 2
[3,] 1 2
> t(matrix)
    [,1] [,2] [,3]
[1,]
[2,] 2 2 2
```

Often you need to select only some column/s or rows of a given object to perform operations:

select row 6 of the object df

```
> df[6,]
  colname 1 colname 2
                 12
select column 2 of the object df
> df[,2]
```

[1] 9 3 25 4 26 12 20 11

Often you need to select only some column/s or rows of a given object to perform operations:

### object[rows, columns]

- select from row 6 to row 8 of the object df

```
> df[6:8,]
  colname_1 colname_2
6          A          12
7          B          20
8          B          11
```

- the same way can be used to remove columns/rows using "-"

#### Remove row 2

Often you need to select only some column/s or rows of a given object to perform operations:

### object[rows, columns]

non-contiguous columns/rows interval must be provided as a vector!

> df[,c(1,3)]

# rows/columns indexing (dataframes)

Columns in dataframes objects can be selected by their names or created using "\$" symbol

9.0 9.5

A 8.5

A 10.0 A 10.5 B 11.0 B 11.5

# rows/columns indexing (dataframes)

Indexes can be used with logical operators to subset values

# rows/columns/object indexing (lists)

select element in list

```
> my list[[1]]  # positional selection (select the first element)
[1] "A" "A" "B" "B" "A" "A" "B" "B"
> my list[["df"]] # selection byname
 colname_1 colname_2 colname_3
               13
                       8.0
        A 5 8.5
               20 9.0
               21 9.5
4
                9 10.0
               28 10.5
6
                8 11.0
               12 11.5
```

# rows/columns/object indexing (lists)

select column in a dataframe included in a list

```
> my_list[["df"]]$colname_1
[1] "A" "A" "B" "B" "A" "A" "B" "B"
```

select rows (or columns) in element included in a list

# filter and subsetting

! for NOT

Filtering and subsetting are the empowered version of row/columns selection that we explained before, but they use specific functions in specific packages such as:

```
subset()
dplyr::filter()
they are based on logics. The grammar for logics in R is simple:
== for equal (not "="
!= for different
> < major an minor (followed by = to include the limit value)
| for OR (this is the "pipe" character usually is before the 1 in keyboards)
& or && for AND</pre>
```



# filter and subsetting

subset() and dplyr::filter() are similar and useful to select certain values in a
dataframe

```
>library(dplyr)
                                              > filter(df, colname 2 < 13)</pre>
> filter(df, colname_1 == "A")
                                                colname_1 colname_2 colname_3
  colname 1 colname 2 colname 3
                                                                           8.5
1
          Α
                   13
                            8.0
                                                        Α
                                                                         10.0
          Α
                            8.5
                                                                         11.0
                                                        В
                                                                   8
                           10.0
5
          Α
                                                        В
                                                                 12
                                                                         11.5
6
                   28
                           10.5
          Α
                                                     subsetting by number
       subsetting by factor
```

# filter and subsetting

```
>filter(df, colname 1 %in% c("B", "C"))  # reads like: "is contained in.."
colname 1 colname 2 colname 3
                 9.0
2 B 23 9.5
    в 8 11.0
             7 11.5
>filter(df, !colname 1 %in% c("B", "C")) # reads like: "is NOT contained in.."
colname 1 colname 2 colname 3
     A
          13
                 8.0
     A 22 8.5
    A 10 10.0
    A 29 10.5
```

### **Exercise**

Create a dataframe from the following vectors:

```
> n <- 1:10
> sex <- rep(c('male', 'female'), 5)
> age <- c(23, 22, 21, 22, 24, 30, 23, 29, 19, 29)
> weight <- c(72, 90, 120, 80, 75, 65, 91, 58, 78, 50)
> height <- c(171, 185, 210, 170, 189, 150, 168, 165, 188, 143)

Question 1: Which is the mean age? [ use mean() function ]</pre>
Question 2: Which is the minimum male weight? [ use function min() ]
```

### **Exercise: solution**

```
> people<- data.frame(n = n, sex = sex, age = age, weight =
weight, height = height)
Question 1: Which is the mean age? [ use mean() function ]
> mean(people$age)
[1] 24.2
Question 2: Which is the minimum male weight?
> min(filter(people, sex=="male")$weight)
[1] 72
                                                  NOTE: in R problems can
                                                  have multiple solutions.
> min(people[people$sex == 'male',]$weight)
                                                  The best one is the
[11 72]
                                                  shortest one (less code
                                                  lines)
```

## Save your session

- \*.R files # your main script (often very light text file data)
- \*.RData files # your environment (all the objects you saved, often very big files (>1Gb)
- You can easily save \*.R and \*.RData files through the RStudio interface
- you can also manage your environments through the R command line (save you some time)

```
# save your environment
```

> save.image(file = "your path/your environment name.RData")

### Save your session

```
# load a *.Data file in your current session
> load(file = "your_path/your_envirnoment_name.RData")
# make searchable a stored environment without loading it
> attach(file = "your_path/your_envirnoment_name.RData")
Note: don't forget to detach() your environment!
```