

# Introduction to R

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## Workshop on Data Visualization in R

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NBIS, SciLifeLab





Only edit title, subtitle & author above this ---





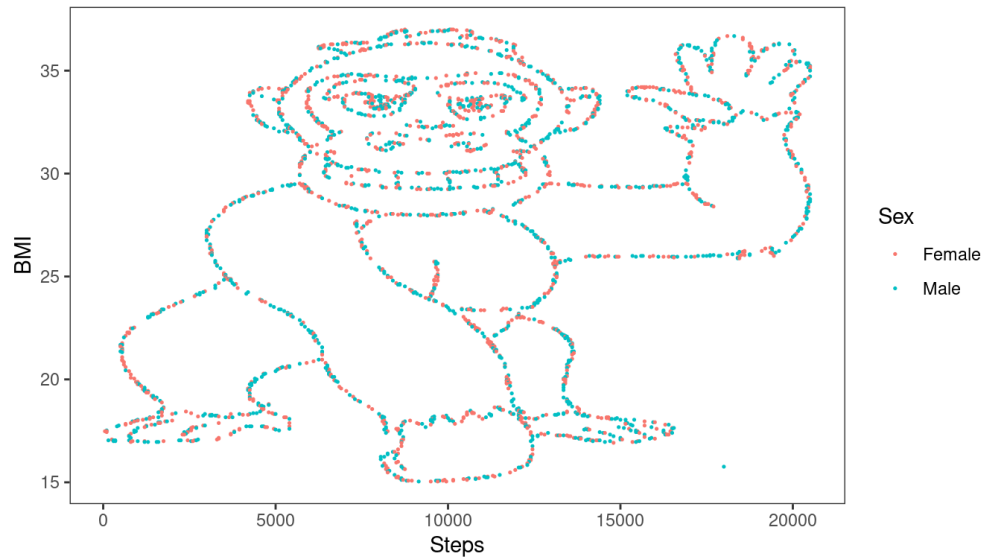
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# Contents

- Course and webpage
- Overview of R
- Data formats
- Data frames
- Important functions
- Tips



# Quick checkups



If you don't recognize the correlation you see in the figure above, I would highly recommend you to read the following paper ;)

**Yanai, I., Lercher, M. A hypothesis is a liability. *Genome Biol* 21, 231 (2020).**

- Coffe breaks (15 minutes is fine?)
- Webpage structure
- Plots from drop-down
- Times mentioned in schedule are **super** arbitrary

# R

- Derived from a statistical programming language called S
- You can write your own functions
- Powerful and flexible.
- Available for all platforms
- **GUI** with Rstudio
- **RMarkdown**: Embedding codes and results together



# Data Formats

- Wide format

	Sample_1	Sample_2	Sample_3	Sample_4
ENSG000000000003	321	303	204	492
ENSG000000000005	0	0	0	0
ENSG000000000419	696	660	472	951
ENSG000000000457	59	54	44	109
ENSG000000000460	399	405	236	445
ENSG000000000938	0	0	0	0

- familiarity
- conveniency
- you see more data

# Data Formats

- Long format

Sample_ID	Gene	count
Sample_1	ENSG00000000003	321
Sample_1	ENSG00000000005	0
Sample_1	ENSG00000000419	696
Sample_1	ENSG00000000457	59
Sample_1	ENSG00000000460	399
Sample_1	ENSG00000000938	0

Sample_ID	Sample_Name	Time	Replicate	Cell	Gene	count
Sample_1	t0_A	t0	A	A431	ENSG00000000003	321
Sample_1	t0_A	t0	A	A431	ENSG00000000005	0
Sample_1	t0_A	t0	A	A431	ENSG00000000419	696
Sample_1	t0_A	t0	A	A431	ENSG00000000457	59
Sample_1	t0_A	t0	A	A431	ENSG00000000460	399
Sample_1	t0_A	t0	A	A431	ENSG00000000938	0

# Data Formats

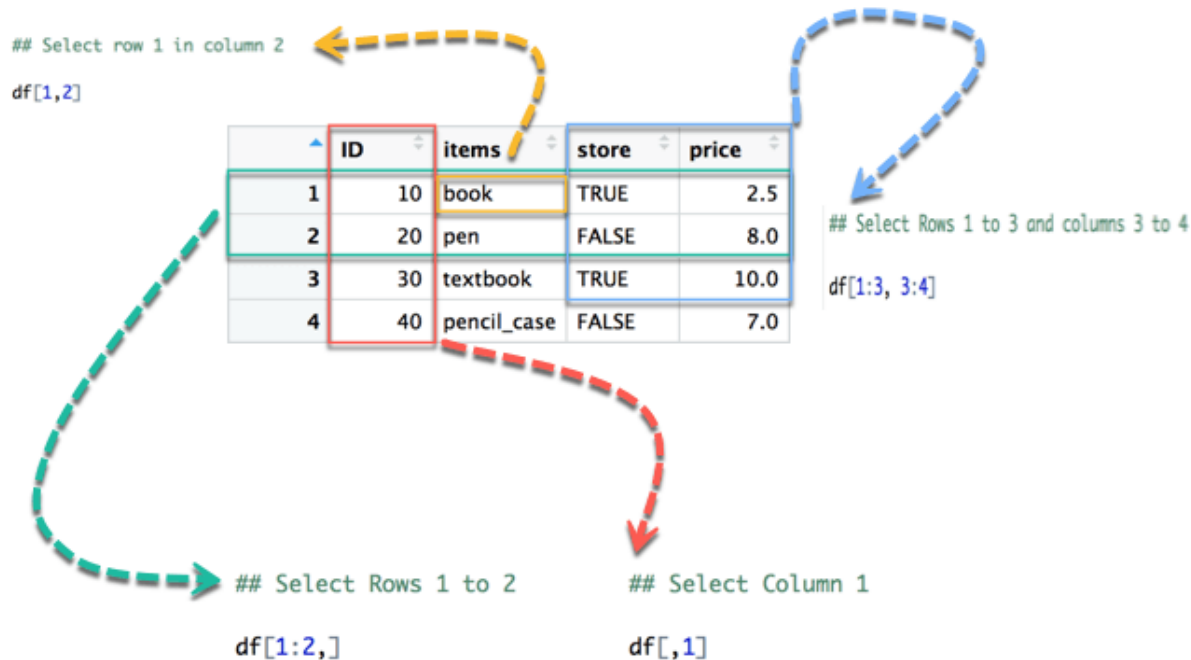
- Long format

Sample_ID	Sample_Name	Time	Replicate	Cell	Gene	count
Sample_1	t0_A	t0	A	A431	ENSG000000000003	321
Sample_1	t0_A	t0	A	A431	ENSG000000000005	0
Sample_1	t0_A	t0	A	A431	ENSG000000000419	696
Sample_1	t0_A	t0	A	A431	ENSG000000000457	59
Sample_1	t0_A	t0	A	A431	ENSG000000000460	399
Sample_1	t0_A	t0	A	A431	ENSG000000000938	0

- easier to add data to the existing
- Most databases store and maintain in long-formats due to its efficiency
- R tools like **ggplot** require data in long format.

# Data Frames

- Let us take a quick look into `data.frame` in `R`:



- imported files are usually in `data.frame`
- Structured matrix with `row.names` and `colnames`
- Probably most used `data.type` in Biology!

# Vectors

```
n <- c(2,3,4,2,1,2,4,5,10,11,8,9)
print(n)
```

```
## [1] 2 3 4 2 1 2 4 5 10 11 8 9
```

```
z <- n +3
print(z)
```

```
## [1] 5 6 7 5 4 5 7 8 13 14 11 12
```

```
z <- n +3
mean(z)
```

```
## [1] 8.083333
```

```
s <- c("I", "love", "Batman")
print(s)
```

```
## [1] "I"      "love"   "Batman"
```

# Vector types

- `int` stands for *integers*
- `dbl` stands for *doubles* or real numbers
- `chr` stands for *character* vectors or strings
- `dtm` stands for *date and time*,
- `lgl` stands for *logical* with just TRUE or FALSE
- `fctr` stands for *factors* which R uses to state categorical variables.
- `date` stands for *dates*

You can find what kind of vectors you have or imported by using the function `class()`





# Thank you. Questions?

R version 4.1.3 (2022-03-10)

Platform: x86\_64-pc-linux-gnu (64-bit)

OS: Ubuntu 18.04.6 LTS

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Built on : 🏠 22-Feb-2023 at 🕒 16:53:55

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