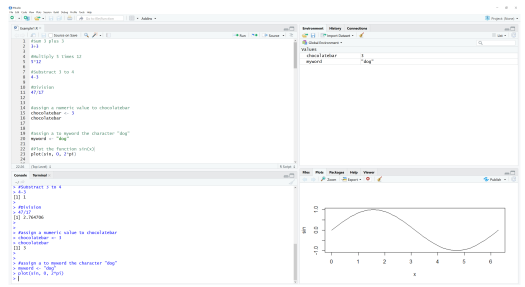
1 Introduction to basic R concepts

- R is a programming language for statistics very popular in biology and bioinformatics.
- It is easy to use, easy to learn, and it is very likely that everything you need has already been coded by someone else.
- It can be used to do statistics at all levels: from simple things such as a t-test or a one-way ANOVA, to advanced statistical modelling such as generalized linear models.
- It is an excellent tool for data visualization. For example Principal Component Analysis (PCA) is widely used to represent bulk RNA-seq data, while t-stochastic neighbourhood embedding (tSNE) is used to represent single cell RNA-seq data.
- During the course we will use Rstudio to code in R.



- The top left panel shows the R script, there is where you write your code.
- The bottom left panel shows the console, there is where things are run.
- The top right panel is the environment, were you can see your stored variables.
- The plots and the help tab are in the lower right panel.
- To create a new R script click on file, New file and Create R script.

- You can save your R script by clicking file and save.
- You can load any saved R script by clicking file and Open file.
- Basic operations:

The input 3+3 gives the output 6,

The input 5*12 gives the output 60,

The input 35/14 gives the output 2.5.

- variables are used to store data. Some basic objects are
 - numerics are used to store numbers.
 chocolate <- 3 stores 3 in the variable chocolate
 soda<-7 stores 7 in the variable soda
 chocolate+soda gives 10.

WARNING: chocolate+Soda gives an error message. R is case sensitive.

- Inf represents a division by 0. 1/0 = Inf. NaN represents not a number, for example 0/0.
- A word writen in quotation "Hello" is readed as a character.word <- "dog"
- Logical objects contain either TRUE or FALSE.
 3<4 will give as an output TRUE.
 7<(4+2) will give as an output FALSE.
- TRUE and FALSE can be used for numerical operations. TRUE has the value 1, FALSE has the value 0. Therefore, TRUE+TRUE+FALSE is 2.
- The type of an object can be checked with the function class()
- A function is a set of actions to perform a task. It requires an input and it produces an output. The structure is NameOfTheFunction(input).
 - mean(c(1,3,8)) gives 4.
 - Mean(c(1,3,8)) gives an error message because R is case sensitive.
 - plot(x=c(1,2,3,4), y=c(1,4,9,16)) plots points at coordinates the coordinates (1,1), (2,4), (3,9) and (4,16).
 - plot(x=c(1,2,3,4), y=c(1,4,9,16), "l") makes the same plot as before, but connecting the points with lines.
- Packages contain one or more functions and, after loading them, those functions are available to use.

- A package is installed with the function install.packages() install.packages("ggplot2") installs the package ggplot2.
- The functions of a package are loaded into your R session with the function library
 - library(ggplot2) loads the package ggplot2.
- You can read the help by typing? in front of a function. ?plot will open the help of the function plot.