



# Reading (and writing) data in R

R Foundations for Life Scientists

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# Reading data



- Reading data is one of the most consuming and most cumbersome aspects of bioinformatics...
- R provides a number of ways to read and write data stored on different media (file, database, url, twitter, Facebook, etc.) and in different formats.
- Package `foreign` contains a number of functions to import less common data formats.

# Reading tables

Most often, we will use the `read.table()` function. It is really, really flexible and nice way to read your data into a data.frame structure with rows corresponding to observations and columns to particular variables.

The function is declared in the following way:

```
read.table(file, header = FALSE, sep = "", quote = "\"'",  
  dec = ".", numerals = c("allow.loss", "warn.loss", "no.loss"),  
  row.names, col.names, as.is = !stringsAsFactors,  
  na.strings = "NA", colClasses = NA, nrows = -1,  
  skip = 0, check.names = TRUE, fill = !blank.lines.skip,  
  strip.white = FALSE, blank.lines.skip = TRUE,  
  comment.char = "#",  
  allowEscapes = FALSE, flush = FALSE,  
  stringsAsFactors = default.stringsAsFactors(),  
  fileEncoding = "", encoding = "unknown", text, skipNul = FALSE)*
```

## `read.table` parameters

You can read more about the *read.table* function on its man page, but the most important arguments are:

- `file` – the path to the file that contains data,
- `header` – a logical indicating whether the first line of the file contains variable names,
- `sep` – a character determining variable delimiter, e.g. comma for csv files,
- `quote` – a character telling R which character surrounds strings,
- `dec` – a character determining the decimal separator,
- `row/col.names` – vectors containing row and column names,
- `na.strings` – a character used for missing data,
- `nrows` – how many rows should be read,
- `skip` – how many rows to skip,
- `as.is` – a vector of logicals or numbers indicating which columns shall not be converted to factors,
- `fill` – add NA to the end of shorter rows,
- `stringsAsFactors` – a logical. Rather self explanatory.

## `read.table` and its siblings



The `read.table` function has some siblings, functions with particular arguments pre-set to a specific value to spare some time:

- `read.csv()` and `read.csv2()` with comma and semicolon as default `sep` and dot and comma as `dec` respectively,
- `read.delim()` and `read.delim2()` for reading tab-delimited files.

We, however, most often use the canonical `read.table()` as it is the most flexible one.

## read.table — example use



```
tab <- read.table(file = 'data/slide_loading_data/2014-04-07_phenos2.csv',  
                  sep = ' ',  
                  header = T)  
tab[1:5, 1:3]  
class(tab$reg_no)
```

```
##           id           uid      reg_no  
## 1 S11799-2007 S11799/2007_621 S11799/2007  
## 2 S14189-2008 S14189/2008_1237 S14189/2008  
## 3 S15108-2008 S15108/2008_1224 S15108/2008  
## 4 S15299-2007 S15299/2007_515 S15299/2007  
## 5 S16860-2009 S16860/2009_849 S16860/2009  
## [1] "character"
```

# What if you encounter errors?

- StackOverflow,
- Google – just type R and copy the error you got without your variable names,
- open the file – has the header line the same number of columns as the first line?
- in Terminal (on Linux/OsX) you can type some useful commands.

## Useful commands for debugging

- `cat phenos.txt | awk -F';' '{print NF}'` prints the number of words in each row. `-F';'` says that semicolon is the delimiter,
- `head -n 5 phenos.txt` prints the 5 first lines of the file,
- `tail -n 5 phenos.txt` prints the 5 last lines of the file,
- `head -n 5 phenos.txt | tail -n 2` will print lines 4 and 5...
- `wc -l phenos.txt` will print the number of lines in the file
- `head -n 2 phenos.txt > test.txt` will write the first 2 lines to a new file

If it still does not give you a clue — just try to load first line of the file.

If this still did not help, split the file in two equal-size parts. Check which part gives the error. Split this part into halves and check which 1/4 gives the error... It is faster than you think!

## Writing with `write.table()`

`read.table()` has its counterpart, the `write.table()` function (as well as its siblings, like `write.csv()`). You can read more about it in the documentation, let us show some examples:

```
vec <- rnorm(10)
write.table(vec, '') # write to screen
write.table(vec, file = 'vector.txt')
# write to the system clipboard, handy!
write.table(vec, 'clipboard', col.names=F,
            row.names=F)
# or on OsX
clip <- pipe("pbcopy", "w")
write.table(vec, file=clip)
close(clip)
# To use in a spreadsheet
write.csv(vec, file = 'spreadsheet.csv')
```



# Writing big data

- HINT: `write.table()` is rather slow on big data – it checks types for every column and row and does separate formatting to each. If your data consists of only one type of data, convert it to a matrix using `as.matrix` before you write it!
- You may want to use function `scan()` that reads files as vectors. The content does not have to be in the tabular form. You can also use scan to read data from keyboard: `typed.data <- scan()`
- If data are written as fixed-width fields, use the `read.fwf()` function.
- Also check out the `readLines()` function that enables you to read data from any stream.

# Read data in xls/xlsx and Matlab



```
library(readxl)
data <- readxl::read_xlsx('myfile.xlsx')
```

```
library(R.matlab)
data <- R.matlab::readMat("mydata.mat")
```

# Working with remote data

```
url <- 'https://en.wikipedia.org/wiki/List_of_countries_by_average_wage'  
conn <- url(url, 'r')  
raw.data <- readLines(conn)  
raw.data[1:3]
```

But data is often tabularized...

```
library(rvest)  
html <- read_html(url)  
tables <- html_nodes(html, 'table')  
data <- html_table(tables[4])[[1]]  
data[1:5, ]
```

# Working with databases

It is also relatively easy to work with different databases. We will focus on MySQL and present only one example that uses the *RMySQL* package (check also *RODBC* and *RPostgreSQL*).

```
library(RMySQL)
db.conn <- dbConnect(MySQL(), user='me',
                     password='qwerty123',
                     dbname='genes',
                     host='127.0.0.237')
query <- dbSendQuery(db.conn, 'SELECT * FROM table7')
data <- fetch(query, n = - 1)
```

See you at the next lecture!

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