Simple data mining in R with data.table and ggplot2

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Reading CSV (or any text tabulated) data

data.table is a package that extends the data.frame object of base R. It comes with its own functions to read (fread) and write (fwrite) tabulated data. Those replacement functions for read.table and write.table are incredibly easy to use and time/memory efficient.

```
library(data.table)
data <- fread(input = "student2/gravi.csv")</pre>
head(data)
        Plante Position NPA time
                                     angle
## 1: Plante 1
                 Courbe Avec
                                 0 168.527
## 2: Plante 2
                                 0 173.283
                 Courbe Avec
## 3: Plante 3
                 Courbe Avec
                                 0 177.797
## 4: Plante 4
                                 0 169.992
                 Courbe Avec
## 5: Plante 5
                 Courbe Avec
                                 0 175.236
## 6: Plante 6
                                 0 177.780
                 Courbe Avec
```

fread is able (most of the time) to automatically identify the type and the layout of your data (separators, headers, ...).

Filtering and manipulating data

The data.table object comes with built-in aggregative capabilities (via the by keyword). A picture is worth a thousand words...

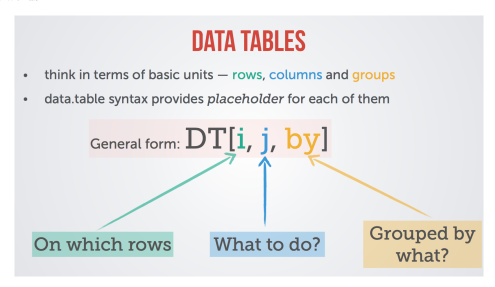


Figure 1: The data.table general form (from https://github.com/Rdatatable/data.table/wiki)

Row filtering

The basic row filtering is similar to the data frame method, except that column names can be used as variables.

```
# The data.frame version
# head(data[data$Position == "Courbe",])

# The data.table version
head(data[Position == "Courbe"])
```

```
Plante Position NPA time
                                     angle
                 Courbe Avec
## 1: Plante 1
                                0 168.527
## 2: Plante 2
                 Courbe Avec
                                0 173.283
## 3: Plante 3
                 Courbe Avec
                                0 177.797
## 4: Plante 4
                 Courbe Avec
                                0 169.992
## 5: Plante 5
                 Courbe Avec
                                0 175.236
## 6: Plante 6
                 Courbe Avec
                                0 177.780
```

Note that with data.table the ',' delimiting the x,y (or i,j) dimensions of the table can be omitted when j is empty (meaning you want to return all the columns without further manipulation).

Data manipulation

At the j placeholder, direct manipulation of data can be perform by using column names as variables.

```
data[Position == "Courbe", mean(angle)]
```

```
## [1] 162.6024
```

The j placeholder accepts a list of arguments, making possible multiple and simultaneous data manipulations.

```
# data[Position == "Courbe", list(mean(angle), sd(angle))]
# The same with direct renaming of the new columns
data[Position == "Courbe", list(mean=mean(angle), sd=sd(angle))]
```

```
## mean sd
## 1: 162.6024 19.63851
```

Data grouping

The third placeholder, by, is used to aggregate data before the manipulation by the j placeholder is performed. by accepts as argument a **list** of one or more column names.

```
data[Position == "Courbe", list(mean=mean(angle), sd=sd(angle)), by=list(NPA, time)]
```

```
## 1: Avec 0 175.0569 4.272599
## 2: Sans 0 172.7916 4.996510
## 3: Avec 1 164.8701 10.752290
## 4: Sans 1 137.6909 23.009854
```

Go-beyond example

The dataset used in the previous examples was made by students who measured the response of *Arabidopsis* thaliana plantlets to a gravitropic stimulus: the plants were either submitted to the gravitropic stimulus

(Position == "Courbe") or not (Position == "Debout") and the role of the phytohormone auxin in this response was evaluated by submitting some of the plants to an auxin transport inhibitor (NPA, NPA == "Avec" vs NPA == "Sans"). The quantification of the response was made by measuring the re-orientation of the hypocotyl one day after the onset of the stimulus, expressed as the difference between the angle (the direction of the growth) measured after (day 1, time == 1) and before (day 0, time == 0) the stimulus. The question was to calculate the mean angle and the standard deviation for each group of experimental conditions.

Here is the way to the one-liner data.table solution.

```
1. For each plantlet in each condition, calculate the 'angle difference' between day 1 and day 0
step1 <- data[, list(diff=angle[time == 1]-angle[time == 0]),</pre>
               by=list(Position, NPA, Plante)]
head(step1)
##
      Position NPA
                       Plante
                                  diff
## 1:
        Courbe Avec Plante 1 -13.452
## 2:
        Courbe Avec Plante 2 -26.212
        Courbe Avec Plante 3 -0.741
## 3:
## 4:
        Courbe Avec Plante 4
                                 1.363
## 5:
        Courbe Avec Plante 5
                                 1.803
## 6:
        Courbe Avec Plante 6
                                 1.050
  2. Calculate the mean and standard deviation for each (Position, NPA) combination of this new data table
step2 <- step1[, list(mean=mean(diff), sd=sd(diff)), by=list(Position, NPA)]</pre>
step2
      Position NPA
##
                         mean
## 1:
        Courbe Avec -10.1868 10.518551
## 2:
        Debout Avec -0.1395 7.567384
## 3:
        Courbe Sans -35.1007 25.521659
## 4:
        Debout Sans -4.3573 13.185171
Both steps can be combined in one line:
my_summary <- data[, list(diff=angle[time == 1]-angle[time == 0]),</pre>
                    by=list(Position, NPA, Plante)][, list(mean=mean(diff), sd=sd(diff)),
                                                       by=list(Position, NPA)]
```

and eventually produce the figure...

```
library(ggplot2)
# Main parameters of the plot: What are my x ? My y ?
# On which parameter do I distinguish my data series ?

g <- ggplot(data = my_summary, mapping = aes(x = Position, y = mean, fill = NPA))

# A 'basic' barplot + error bars
g.basic <- g +
    geom_errorbar(width=.25, aes(ymin=mean-sd, ymax=mean), position = position_dodge(0.9)) +
    geom_bar(stat = "identity", position = position_dodge(), color = "black")

g.basic</pre>
```

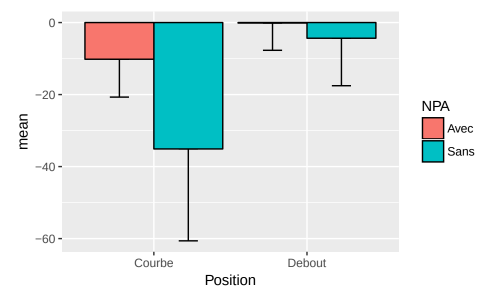


Figure 2: My basic plot.

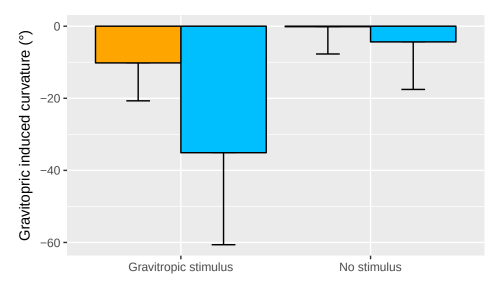


Figure 3: My improved plot.