

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")
head(data)
```

```
##      Plante Position  NPA time   angle
## 1: Plante 1   Courbe Avec    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
```

`fread` is able (most of the time) to automatically identify the type and the layout of your data (separators, headers, ...). If your data are encoded with a decimal separator other than the `.`, it is necessary to set the `dec` argument of `fread`: `data <- fread(input = "student2/gravi.csv", dec=",")`.

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...

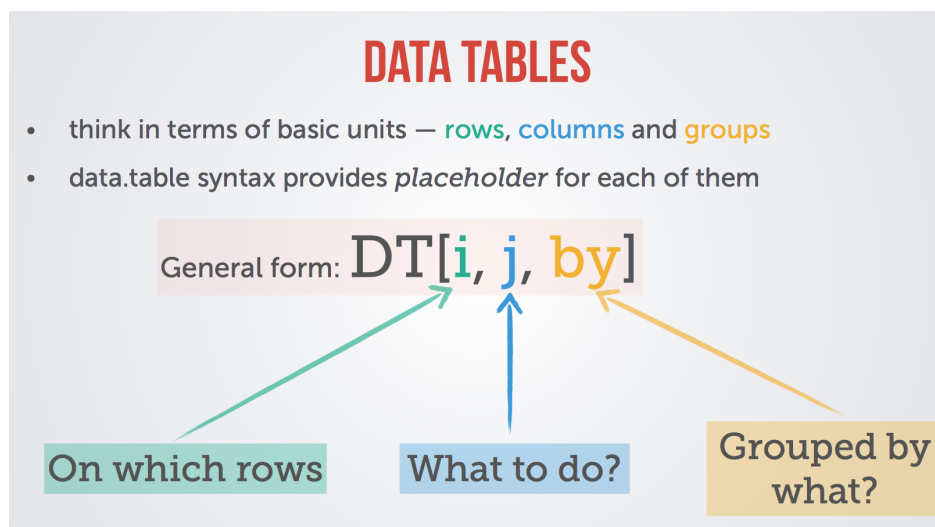


Figure 1: The `data.table` general form (from <https://github.com/Rdatatable/data.table/wiki>)

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
## 1: Plante 1   Courbe Avec    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 performed 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
```

‘On place’ update of `data.table` objects

The `:=` operator can be used at the *j* placeholder to add, remove and update columns by reference. For example, if you want to transform, in place, the angle values from degrees to radians, just use :

```
# Update the angle column with radian values
head(data[, angle := angle * (pi/180)])
```

```
##      Plante Position  NPA time   angle
## 1: Plante 1   Courbe Avec    0 2.941351
## 2: Plante 2   Courbe Avec    0 3.024359
## 3: Plante 3   Courbe Avec    0 3.103143
## 4: Plante 4   Courbe Avec    0 2.966920
## 5: Plante 5   Courbe Avec    0 3.058445
## 6: Plante 6   Courbe Avec    0 3.102846
```

Note that, by using the `:=` operator, you modify your original `data.table` object... even if you do not use explicitly the `<-` assignment operator, like in `data[, angle := angle * (pi/180)]`.

This operator can also be used to add a new column or remove an existing column:

```
# reload the original file since previous commands modified the table
data <- fread(input = "student2/gravi.csv")
# add a column with radian values
data[, rad := angle * (pi/180)]
# remove the angle column
data[, angle := NULL]
```

or in one operation:

```
# reload the original file since previous commands modified the table
data <- fread(input = "student2/gravi.csv")
head(data[, c("rad", "angle") := list(angle * (pi/180), NULL)])
```

```
##      Plante Position  NPA time      rad
## 1: Plante 1   Courbe Avec    0 2.941351
## 2: Plante 2   Courbe Avec    0 3.024359
## 3: Plante 3   Courbe Avec    0 3.103143
## 4: Plante 4   Courbe Avec    0 2.966920
## 5: Plante 5   Courbe Avec    0 3.058445
## 6: Plante 6   Courbe Avec    0 3.102846
```

```
# equivalent to
# data[, `:=` (rad = angle * (pi/180), angle = NULL)]
```

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.

```
# reload the original file since previous commands modified the table
data <- fread(input = "student2/gravi.csv")
data[Position == "Courbe",
     list(mean=mean(angle), sd=sd(angle)),
     by=list(NPA, time)]
```

```
##      NPA time      mean      sd
## 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]),  
               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  
## 3: Courbe Avec Plante 3  -0.741  
## 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)]  
step2
```

```
##   Position  NPA      mean      sd  
## 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]),  
                   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  
# (the 'fill' argument is used to produced grouped bars on the NPA data)  
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  
  
# With the same data, rather than grouped bars, NPA data is used  
# to separate the bar plot in 2 facets  
g.basic <- g +  
  geom_errorbar(width=.25, aes(ymin=mean-sd, ymax=mean), position = position_dodge(0.9)) +
```

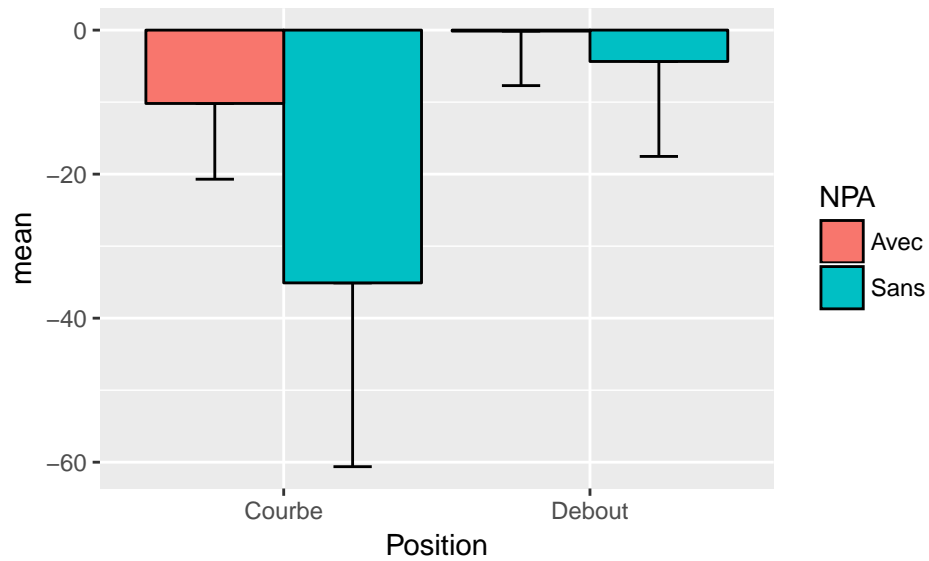


Figure 2: My basic plot.

```
geom_bar(stat = "identity", position = position_dodge(), color = "black") +
facet_grid(. ~ NPA, labeller = labeller(NPA = c(Avec="+NPA", Sans="-NPA")))
```

g.basic

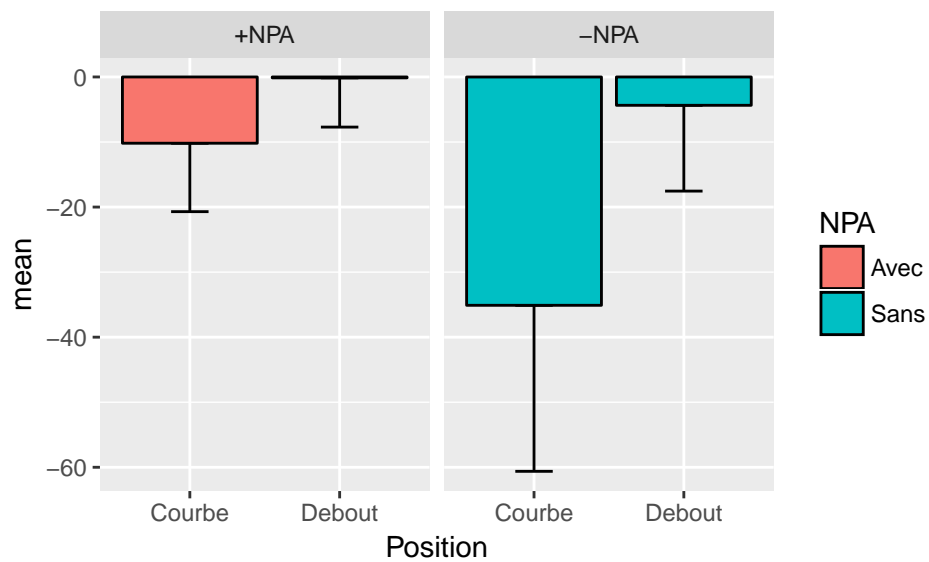


Figure 3: My basic faceted plot.

```
# An improved plot
g.improved <- g.basic +
  scale_fill_manual(values = c("orange", "deepskyblue")) + # changing the colors
  scale_x_discrete(labels=c("Gravitropic\nstimulus",
                           "No stimulus")) + # changing the x tick labels
  ylab("Gravitropic induced curvature (°)") + # The y axis title
  xlab("") + # The x axis title
```

```
guides(fill=FALSE) # Hide the legend
```

```
g.improved
```

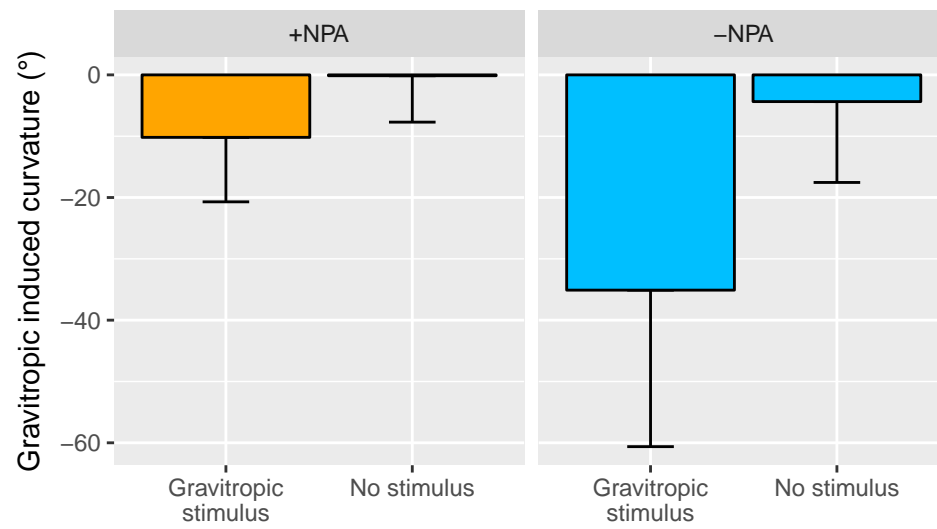


Figure 4: My improved plot.