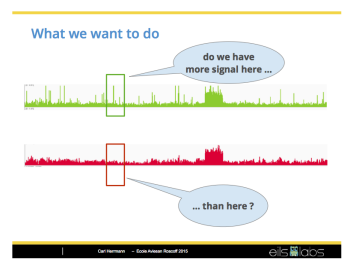
Practical: basic stats for peak calling

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Peak-calling: Carl's question



Defining the data directory

Open a connection to the lab web browser.

We will first define the URL from which the data can be downloaded, by concatenating the URL fo the course with the path to our dataset.

To concatenate paths, it is *recommended* to use the $\bf R$ command file.path().

```
url.course <- "http://jvanheld.github.io/stats_avec_RStudio
url.data <- file.path(url.course, "practicals", "02_peak-categories")</pre>
```

Loading a data table

R enables to download data directly from the Web. Load counts per window in chip sample.

```
## Define URL of the ChIP file
chip.bedg.file <- file.path(url.data, "FNR_200bp.bedg")

## Load the file content in an R data.frame
chip.bedg <- read.table(chip.bedg.file)

## Set column names
names(chip.bedg) <- c("chrom", "start", "end", "counts")</pre>
```

Exploring a data frame: dim()

Before anything else, let us inspect the size of the data frame, in order to check that it was properly lodaded.

```
dim(chip.bedg)
```

```
## [1] 23199
```

Checking the n first rows: head()

The function head() displays the first rows of a table.

```
head(chip.bedg, n = 5)
```

```
##
                             chrom start end counts
   1 gi|49175990|ref|NC_000913.2|
                                          200
                                                 1594
## 2 gi|49175990|ref|NC_000913.2|
                                     200
                                          400
                                                  834
## 3 gi|49175990|ref|NC_000913.2|
                                     400
                                          600
                                                  222
## 4 gi|49175990|ref|NC_000913.2|
                                     600
                                          800
                                                  172
## 5 gi|49175990|ref|NC_000913.2|
                                     800 1000
                                                  123
```

Checking the n last rows: tail()

The function tail() displays the first rows of a table.

```
tail(chip.bedg, n = 5)
```

```
## 23195 gi|49175990|ref|NC_000913.2| 4638800 4639000 19
## 23196 gi|49175990|ref|NC_000913.2| 4639000 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200 4639200
```

Viewing a table

The function View() displays the full table in a user-friendly mode.

View(chip.bedg)

Selecting arbitrary rows

```
chip.bedg[100:105,]
```

```
## 100 gi|49175990|ref|NC_000913.2| 19800 20000 21

## 101 gi|49175990|ref|NC_000913.2| 20000 20200 0

## 102 gi|49175990|ref|NC_000913.2| 20200 20400 0

## 103 gi|49175990|ref|NC_000913.2| 20400 20600 108

## 104 gi|49175990|ref|NC_000913.2| 20600 20800 229

## 105 gi|49175990|ref|NC_000913.2| 20800 21000 245
```

Selecting arbitrary columns

```
chip.bedg[100:105, 2]
## [1] 19800 20000 20200 20400 20600 20800
chip.bedg[100:105, "start"]
## [1] 19800 20000 20200 20400 20600 20800
chip.bedg[100:105, c("start", "counts")]
## start counts
## 100 19800
                21
## 101 20000 0
## 102 20200
## 103 20400 108
## 104 20600
               229
                                   4 D > 4 B > 4 B > 4 B > 9 Q P
```

Adding columns

We can add columns with the result of computations from other columns.

```
chip.bedg$midpos <- (chip.bedg$start + chip.bedg$end)/2
head(chip.bedg)</pre>
```

```
##
                             chrom start
                                           end counts midpos
   1 gi|49175990|ref|NC 000913.2|
                                                          100
                                           200
                                                  1594
   2 gi|49175990|ref|NC 000913.2|
                                           400
                                                   834
                                                          300
                                      200
   3 gi|49175990|ref|NC 000913.2|
                                                   222
                                                          500
                                      400
                                           600
   4 gi|49175990|ref|NC 000913.2|
                                                   172
                                                          700
                                      600
                                           800
## 5 gi|49175990|ref|NC 000913.2|
                                      800
                                          1000
                                                   123
                                                          900
## 6 gi|49175990|ref|NC 000913.2|
                                                   116
                                                         1100
                                     1000 1200
```

Plotting a density profile

We can readily print a plot with the counts per window.

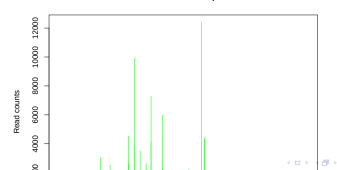
2e+06

```
plot(chip.bedg[, c("midpos", "counts")], type="h")
   12000
   10000
   8000
   0009
   2000
```

Plotting a density profile

Let us improve the plot

FNR ChIP-seq



Exercise: exploring the background

We already loaded the count table for the FNR ChIP counts per window.

The background level will be estimated by loading counts per window in a genomic input sample. These counts are available in the same directory a file named input_200bp.bedg

- 1. Load the counts per window in the input sample (genome sequencing).
- 2. Plot the density profile of the input
- 3. Compare chip-seq and input density profiles
- 4. Compare counts per window between chip-seq and input

Solution: loading the input counts per window

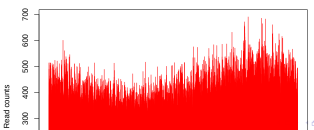
```
## Define URL of the input file
input.bedg.file <- file.path(url.data, "input_200bp.bedg")

## Load the file content in an R data.frame
input.bedg <- read.table(input.bedg.file)

## Set column names
names(input.bedg) <- c("chrom", "start", "end", "counts")</pre>
```

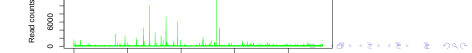
Solution: plotting the input density profile

Background (genomic input)



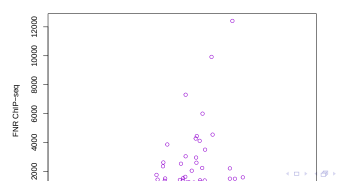
Solution: comparing chip-seq and background density profiles

FNR ChIP-seq



Solution: comparing counts per window between chip-seq and input

Read counts per 200bp window



Solution: comparing counts per window between chip-seq and input

In order to better highight the dynamic range, we can use a log-based representation

```
## Warning in xy.coords(x, y, xlabel, ylabel, log): 377 x v
## from logarithmic plot
## Warning in xy.coords(x, y, xlabel, ylabel, log): 403 y v
## from logarithmic plot
```

```
grid() ## add a grid
```

Questions

- On the ChIP-seq versus input plot, how would you define peaks
- ► Where would you place the limit between peaks and background fluctuations ?

Exercises

- 1. Think about further drawing modes to improve your perception of the differences between signal and background.
- 2. We will formulate (together) a reasoning path to compute a p-value for each peak.