Analysis of a genome annotation table

Probabilities and statistics for biology (STAT1)

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Goal of this practical

During this practical session, you will run the following tasks:

- 1. Handle a table containing annotated features of the yeast genome.
- 2. Select a subset of the data by filtering rows based on a given criterion (annotation type, chromosome, \dots)
- 3. Generate graphics to represent different aspects of the data.
- 4. Compute estimators of central tendency and dispersion.
- 5. Compute a confidence interval around the mean.

Expected report

At the end of the practical you will be asked to submit two documents

- 1. Your **R** code. Each question must be explicitly formulated before presenting the results that answer it and giving an interpretation of these results.
- 2. UA **synthetic report**, which will include a presentation of the main results (figures, descriptive stats, tables) as well as your interpretation of the result.

Expectation for the code

- 1. The code must be **readable and undestandable**: choose variable names that explicitly indicate what they represent.
- 2. The code must be properly documented (the # symbol starts a comment, either at the beginning or in the middle of a line of code).
 - Before each chunk of code, explain what this code is supposed to do, what it serves to.
 - Don't hesitate to occasionally add some comment words to justify the chosen approach.
 - Each time you define a variable, add a comment on the same line to indicate what this variable represents.
- 3. The code must be **portable**: other people should be able to download it and run it on their computer. For this practical, I will systematically test whether your code can run on my computer. hard-coded absolute paths of a file on your machine should thus always be avoided (we will indicate hereafter how to define relative paths relative to the root of your user account).

Expected content for the interpretation report

Your report must be synthetic (1 text page $\max +$ as many figures and table as you wish)

Each question must be explicitly formulated before presenting the results that answer it and then interpreting those results.

Each figure or table must be documented with a legend that allows a naive reader to understand what it represents. The interpretation of the results displayed on a figure or table will be found in the main text (with a reference to the figure or table number).

Historical example: yeast genome

- 1992: publication of the first complete eukaryotic chromosome, the 3rd yeast chromosome.
- 1996: publication of the complete genome.

On the base of the genes of the 3rd chromosome (sample) we can estimate the average size of a yeast gene.

Questions:

- (a) Would the sample mean (chromosome III) be sufficient to predict the population mean (complete genome) ?
 - To answer this question, we will imagine that we came back in 1992, and will use all the genes of chromosome III (considered here as a sample of the genome) to estimate the average size of genes for the whole genome (the "population" of genes").
- (b) Can this sample be described as "simple and independent"?

Analysis of the length of the baker's yeast genes

Tutorial

Before moving to the exercises, we show you here some basic elements about reading, manipulating and writing data tables with R.

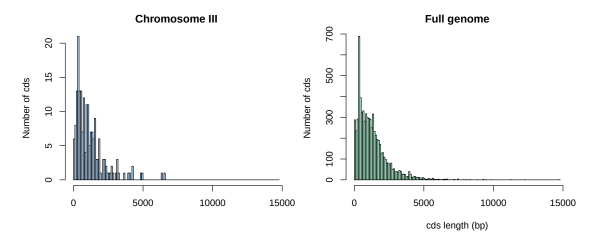


Figure 1: Distribution of cds lengths for Saccharomyces cerevisiae.

The path to the home (manual)

We will create a folder for this tutorial, starting from the root of our account.

First possibility (quick but not very elegant): enter (manually) the path from the root of your account in a variable

dir.home <- /the/path/to/the/home</pre>

- Advantage: fast and convenient
- Disadvantage: not portable, will only work on your computer

The path to the home (automatic)

A more general solution: use the R command Sys.getenv().

- Invoked without parameters, this command lists all environment variables (your system configuration).
- The output can be restricted to a given environment variable, for example Sys.getenv("HOME") returns the path to the root of your account.

Note: equivalent writing with Linux: the tilde symbol ~ also indicates the path to the root of your account.

```
## Identify the home directory
## by getting the environment variable HOME
dir.home <- Sys.getenv("HOME")
print(dir.home)</pre>
```

[1] "/Users/jvanheld"

Creating a folder for the TP

```
## Define a variable containing the path of the results for this tutorial
dir.tuto <- file.path(dir.home, "stat1", "TP2")
print(dir.tuto)</pre>
```

[1] "/Users/jvanheld/stat1/TP2"

```
## Create the directory for this tutorial
dir.create(path = dir.tuto, showWarnings = FALSE, recursive = TRUE)
```

```
## Go to the tutorial directory
setwd(dir.tuto)

## List the files already present in the folder (if any)
list.files()
```

- [1] "3nt_genomic_Saccharomyces_cerevisiae-ovlp-1str.tab"
- [2] "chrom sizes.tsv"
- [3] "Saccharomyces_cerevisiae.R64-1-1.37.gtf.gz"

Downloading the GTF file from EnsemblGenomes

Tips: before downloading the annotation file (GTF) from EnsemblGenomes to our computer, we will check if it is already present (and in this case we do not re-download it).

```
## Define the URL of the annotation file (GTF-formatted)
gtf.URL <- "ftp://ftp.ensemblgenomes.org/pub/release-37/fungi/gtf/saccharomyces_cerevisiae/Saccharomyce

## Define the path where the local copy will be stored
local.GTF <- file.path(dir.tuto, "Saccharomyces_cerevisiae.R64-1-1.37.gtf.gz")

## If the local file file laready exists, skip the download
if (file.exists(local.GTF)) {
   message("GTF file already exists in the tutorial folder: ", local.GTF)
} else {
   ## Download annotation table in GTF format
   download.file(url = gtf.URL, destfile = local.GTF)
}</pre>
```

Loading a data table

R has several types of tabular structures (matrix, data.frame, table).

The most commonly used structure is the data.frame, which consists of an array of values (numeric or strings) whose rows and columns are associated with names.

The function read.table() allows you to read a text file containing a data table, and store the content in a variable.

Several functions derived from read.table() make it easier to read different types of formats:

- read.delim() for files whose columns are delimited by a particular character (usually the tab, represented by "").
- read.csv() for files "comma-separated values".
- 1. Download the following file to your computer:
- Saccharomyces_cerevisiae.R64-1-1.37.gtf
- 2. Load it using the read table function (for this you must replace the path below by that of your computer).

```
## Read a GTF file with yeast genome annotations

## Load the feature table
feature.table <- read.table(
  local.GTF,
  comment.char = "#",
  sep="\t",
  header=FALSE,</pre>
```

Exploring the content of a data table

The first thing to do after loading a data table is to check its dimensions.

```
dim(feature.table) ## Dimensions of the tbale
```

[1] 43028 9

```
nrow(feature.table) ## Number of rows
```

[1] 43028

```
ncol(feature.table) ## Number of columns
```

[1] 9

The display of the complete annotation table would not be very readable, since it contains tens of thousands of lines.

We can display the first lines with the function head().

Note: the last column is particularly heavy (it contains a lot of information). We will see later how to select a subset of the columns to simplify the display.

```
## Display the 5 first rows of the feature table
head(feature.table, n = 5)
```

```
seqname source
                     feature start end score strand frame
1
       ΙV
             SGD
                        gene 1802 2953
2
       ΙV
             SGD transcript
                              1802 2953
3
       IV
             SGD
                        exon
                              1802 2953
4
       ΙV
             SGD
                         CDS
                              1802 2950
                                                          0
                                                          0
5
       ΙV
             SGD start_codon
                             1802 1804
```

```
gene_id YDL248W; transcript_id YDL248W; gene_name cost; gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source 4 gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_biotype is gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source score is gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source score is gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source score is gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source score is gene_id YDL248W; transcript_id YDL248W; exon_number 1; gene_name cost; gene_source score is gene_id YDL248W; exon_number is gene_id YDL248
```

The function tail() displays the last few lines:

```
## Display the 5 last rows of the feature table
tail(feature.table, n = 5)
```

	seqname	source	feature	start	end	score	${\tt strand}$	frame
43024	Mito	SGD	transcript	85554	85709		+	
43025	Mito	SGD	exon	85554	85709		+	
43026	Mito	SGD	CDS	85554	85706		+	0
43027	Mito	SGD	start_codon	85554	85556		+	0
43028	Mito	SGD	stop codon	85707	85709		+	0

```
gene_id Q0297; transcript_id Q0297; gene_sour
43025 gene_id Q0297; transcript_id Q0297; exon_number 1; gene_source SGD; gene_bioty
43026 gene_id Q0297; transcript_id Q0297; exon_number 1; gene_source SGD; gene_biotype protein_coding;
43027 gene_id Q0297; transcript_id Q0297; exon_number 1; gene_sour
43028 gene_id Q0297; transcript_id Q0297; exon_number 1; gene_sour
```

If you are using the **RStudio** environment, you can display the table in a dynamic viewer pane with the function View().

```
## In RStudio, display the table in a separate tab
View(feature.table)
```

Selection of subsets from a table

Selection of a line specified by its index.

feature.table[12,]

12 gene_id YDL247W-A; transcript_id YDL247W-A; exon_number 1; gene_source SGD; gene_biotype protein_cod Selection of a column specified by its index (display of the first values only).

head(feature.table[,3])

```
[1] gene transcript exon CDS start_codon stop_codon Levels: CDS exon gene start_codon stop_codon transcript
```

Selection of a cell by combining row and column indices.

feature.table[12, 3]

[1] stop_codon

Levels: CDS exon gene start_codon stop_codon transcript

Selection of a column and/or row set.

feature.table[100:105, 1:6]

	seqname	source	feature	start	end	score
100	IV	SGD	CDS	34240	36477	
101	IV	SGD	start_codon	36475	36477	
102	IV	SGD	stop_codon	34237	34239	
103	IV	SGD	gene	36797	38173	
104	IV	SGD	transcript	36797	38173	•
105	IV	SGD	exon	36797	38173	

Selection of specific columns (here, the genomic coordinates of each feature): chromosome, beginning, end, strand.

feature.table[100:105, c(1,4,5,7)]

strand	end	start	seqname	
_	36477	34240	IV	100
_	36477	36475	IV	101
-	34239	34237	IV	102
+	38173	36797	IV	103
+	38173	36797	IV	104
+	38173	36797	IV	105

Select a column based on its name.

```
head(feature.table$start, n=100)
  [1] 1802 1802 1802 1802 1802 2951
                                           3762 3762
                                                     3762
                                                            3762
                                                                   3762
             5985 5985 5985
 [12]
                              5985
                                     5985
                                           7812
                                                 8683
                                                       8683
                                                            8683
 [23] 9754 8683 11657 11657 11657 11660 13358 11657 16204 16204 16204
 [34] 16204 16204 17224 17577 17577 17577 17580 18564 17577 18959 18959
 [45] 18959 18959 18959 19310 20635 20635 20635 20635 20635 21004 22471
 [56] 22471 22471 22474 22606 22471 22823 22823 22823 22823 22823 25874
 [67] 26403 26403 26403 26406 28773 26403 28985 28985 28985 28988 30452
 [78] 28985 30657 30657 30657 30657 30657 31827 32296 32296 32296 32296
 [89] 32296 33232 33415 33415 33415 33418 33916 33415 34237 34237 34237
[100] 34240
head(feature.table$feature, n=20)
```

```
[1] gene transcript exon CDS start_codon
[6] stop_codon gene transcript exon CDS
[11] start_codon stop_codon gene transcript exon
[16] CDS start_codon stop_codon gene transcript
Levels: CDS exon gene start_codon stop_codon transcript
```

Selection of several columns based on their names.

```
## Select the "start" column and print the 100 first results
feature.table[100:106, c("seqname", "start", "end", "strand")]
```

```
seqname start
                     end strand
100
         IV 34240 36477
101
         IV 36475 36477
         IV 34237 34239
102
103
         IV 36797 38173
104
         IV 36797 38173
105
         IV 36797 38173
         IV 36797 38170
106
```

Note: Selection of several columns based on their names. It is also possible to name the rows of a data.frame but the GTF table does not support this. We will see more examples later.

Selection of a subset of rows based on the content of a column

The function subset() allows you to select a subset of the rows of a data.frame based on a condition applied to one or more columns.

We can apply it to select the subset of rows in the annotation table corresponding to coding sequences (CDS).

```
## Select subset of features having "cds" as "feature" attribute
cds <- subset(feature.table, feature=="CDS")
nrow(feature.table) ## Count the number of features</pre>
```

[1] 43028

```
nrow(cds) ## Count the number of cds
```

[1] 7050

Count by value

The function table() allows you to count the occurrences of each value in a vector or array. Some examples of use below.

```
## Count the number of features per chromosome
table(feature.table$seqname)
```

```
I II III IV IX Mito V VI VII VIII X XI XII XIII XIV 759 2912 1210 5374 1567 327 2159 946 3856 2054 2617 2231 3789 3311 2774 XV XVI 3846 3296
```

```
## Count the number of features per type
table(feature.table$feature)
```

```
CDS exon gene start_codon stop_codon transcript 7050 7872 7445 6700 6516 7445
```

Contingency tables can be calculated by counting the number of combinations between 2 vectors (or 2 columns of a table).

```
## Table with two vectors
table(feature.table$feature, feature.table$seqname)
```

```
I II III IV IX Mito
                                       V VI VII VIII
                                                         X XI XII XIII
            122 492 194 895 255
CDS
                                  59 345 151 619
                                                  346 422 361 615
                                                                   544
exon
            137 525 224 961 288
                                  94 400 180 710
                                                  373 480 404 698
                                                                    610
gene
            132 494 213 914 274
                                  62 383 167 676
                                                  349 458 388 658
                                                                    573
start codon 119 464 185 853 243
                                  28 328 143 593
                                                  325 406 348 586
                                                                    514
stop_codon 117 443 181 837 233
                                  22 320 138 582
                                                  312 393 342 574
                                                                    497
transcript
            132 494 213 914 274
                                  62 383 167 676
                                                  349 458 388 658
            XIV XV XVI
CDS
            458 623 549
            500 689 599
exon
            475 665 564
gene
start_codon 438 607 520
stop_codon
            428 597 500
            475 665 564
transcript
```

```
## Same result with a 2-column data frame
table(feature.table[, c("feature", "seqname")])
```

```
seqname
feature
                I II III IV IX Mito
                                         V VI VII VIII
                                                          X XI XII XIII
  CDS
              122 492 194 895 255
                                    59 345 151 619
                                                    346 422 361 615
                                                                     544
  exon
              137 525 224 961 288
                                    94 400 180 710
                                                    373 480 404 698
                                                                     610
  gene
              132 494 213 914 274
                                    62 383 167 676
                                                    349 458 388 658
  start_codon 119 464 185 853 243
                                    28 328 143 593
                                                    325 406 348 586
                                                                     514
  stop_codon 117 443 181 837 233
                                    22 320 138 582
                                                    312 393 342 574
                                                                     497
                                    62 383 167 676 349 458 388 658 573
  transcript
             132 494 213 914 274
             seqname
feature
              XIV XV XVI
  CDS
              458 623 549
```

```
exon 500 689 599 gene 475 665 564 start_codon 438 607 520 stop_codon 428 597 500 transcript 475 665 564
```

Exercises

1. GTF format specifications

Read the GTF format specifications.

- Ensembl (http://www.ensembl.org/info/website/upload/gff.html)
- UCSC (https://genome.ucsc.edu/FAQ/FAQformat.html#format4)

2. Creating a local folder for the TP

Create a local folder (for example: stat1/TP_yeast from the root of your account). We suggest you to use the following functions:

- Sys.getenv("HOME") (Linux and Mac OS X), to get the root of your user account;
- file.path() to build a path;
- dir.create() to create the folder for the TP. Read carefully the options of this function with help(dir.create)

3. Locating the annotation file

Locate the yeast genome annotation file in GTF format in this local folder.

- Site Ensembl Fungi: http://fungi.ensembl.org/
- Click "Downloads" to access the ftp website
- In the search box, type "saccharomyces cerevisiae" and follow the link "GTF"
- Copy the address (URL) of the file Saccharomyces cerevisiae.R64-1-1.37.gtf.gz

4. Downloading a file from an ftp website

Suggested functions:

• download.file() (read the help to know the arguments)

5. Loading a data table in R

Write a script that loads the data table into a variable named feature.table, using the function R read.delim().

Be sure to ignore the comment lines (which start with a character #).

6. Compute the length of coding genes

• Add to the annotation table (feature.table) a column entitled "length" which indicates the length of each annotated genomic feature.

```
## Add a colmn with feature lengths
feature.table[, "length"] <- feature.table[, "end"] - feature.table[, "start"] + 1

## Add a colmn with feature lengths: equivalent result with simpler notation
feature.table$length <- feature.table$end - feature.table$start + 1</pre>
```

• Count the number of rows in the table corresponding to each type of annotation (3rd column of the GTF, "feature").

- fonction table()

~table(feature.table\$feature)

- Select the lines corresponding to coding regions ("CDS")
 - fonction subset()
- Count the number of CDS per chromosome.
 - fonction table()

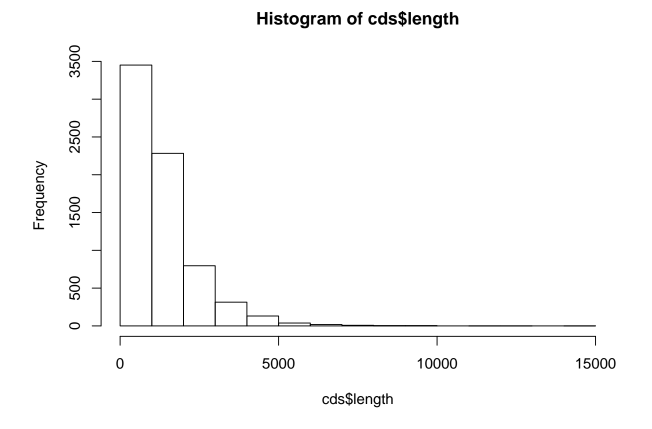
```
Ι
      II
           III
                        IX Mito
                                         VI
                                              VII VIII
                                                           Х
                                                                XΙ
                                                                     XII XIII
122
     492
           194
                 895
                       255
                                                               361
                                                                     615
                                                                                458
                             59
                                  345
                                        151
                                              619
                                                   346
XV
     XVI
     549
623
```

• Load the chromosome size table chrom_sizes.tsv, and calculate the gene density for each chromosome (number of genes per Mb).

[1] 316617

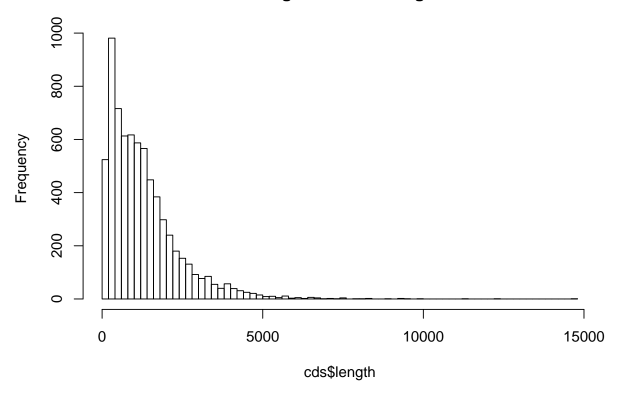
6. Histogram of gene length

By using the function hist(), draw a histogram representing the length distribution of the CDS.

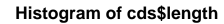


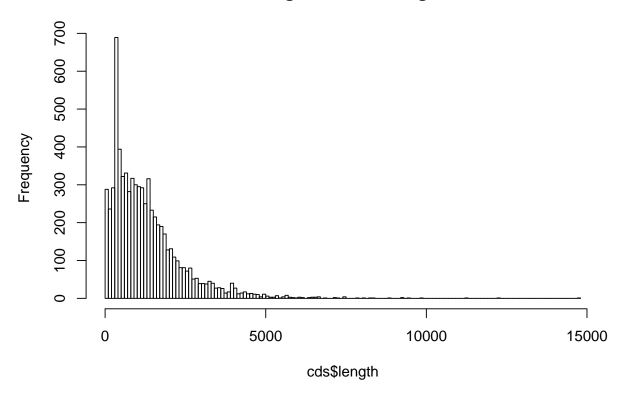
Choose the class intervals in a way that the histogram is informative (neither too large nor too few classes).

Histogram of cds\$length



Retrieve the result of hist() in a variable named cds.length.hist.





Print the result on the screen (print()) and analyze the structure of the variable cds.length.hist (this is a list variable).

Useful functions:

\$breal	KS																			
[1]		0	100	20	00	300	40	00		500	60	00	•	700	80	00		900	100	00
[12]	110	00	1200	130	00	1400	150	00	1	600	170	00	18	300	190	00	2	000	210	00
[23]	220	00	2300	240	00	2500	260	00	2	700	280	00	29	900	300	00	3	100	320	00
[34]	330	00	3400	350	00	3600	370	00	3	800	390	00	40	000	410	00	4	200	430	00
[45]	440	00	4500	460	00	4700	480	00	4	900	500	00	5:	100	520	00	5	300	540	00
[56]	550	00	5600	570	00	5800	590	00	6	000	610	00	62	200	630	00	6	400	650	00
[67]	660	00	6700	680	00	6900	700	00	7	100	720	00	73	300	740	00	7	500	760	00
[78]	770	00	7800	790	00	8000	810	00	8:	200	830	00	84	100	850	00	8	600	870	00
[89]	880	00	8900	900	00	9100	920	00	9	300	940	00	9	500	960	00	9	700	980	00
[100]	990	00 1	0000	1010	00 1	0200	1030	00	10	400	1050	00	106	300	1070	00	10	800	1090	00
[111]	1100	00 1	1100	1120	00 1	1300	1140	00	11	500	1160	00	11	700	1180	00	11	900	1200	00
[122]	1210	00 1	2200	1230	00 1	2400	1250	00	12	600	1270	00	128	300	1290	00	13	000	1310	00
[133]	1320	00 1	3300	1340	00 1	3500	1360	00	13	700	1380	00	139	900	1400	00	14	100	1420	00
[144]	1430	00 1	4400	1450	00 1	4600	1470	00	14	800										
\$count	ts																			
[1]	288	236	292	689	394	322	331	28	32	317	300	29	5 2	292	250	31	6	233	215	194
[18]	190	170	128	131	109	99	81	8	31	72	80	5	51	53	39	3	9	38	45	40
[35]	27	28	26	14	17	40	27	1	.2	14	17	1	2	13	11	1	0	4	11	6
[52]	3	3	7	1	4	8	3		2	1	3		2	0	2		3	3	4	0

[69]	1	0	0	2	1	0	4	0	0	0	1	0	1	0	1	1	0
[86]	0	0	0	1	0	0	0	2	0	1	0	0	0	1	0	0	0
[103]	0	0	0	0	0	0	0	0	0	0	1	0	0	0	0	0	0
[120]	0	0	0	1	0	0	0	0	0	0	0	0	0	0	0	0	0
[137]	0	0	Ο	0	Ο	0	Ο	Ο	0	Ο	Ο	1					

\$density

[1] 4.085106e-04 3.347518e-04 4.141844e-04 9.773050e-04 5.588652e-04 [6] 4.567376e-04 4.695035e-04 4.000000e-04 4.496454e-04 4.255319e-04 [11] 4.184397e-04 4.141844e-04 3.546099e-04 4.482270e-04 3.304965e-04 [16] 3.049645e-04 2.751773e-04 2.695035e-04 2.411348e-04 1.815603e-04 [21] 1.858156e-04 1.546099e-04 1.404255e-04 1.148936e-04 1.148936e-04 [26] 1.021277e-04 1.134752e-04 7.234043e-05 7.517730e-05 5.531915e-05 [31] 5.531915e-05 5.390071e-05 6.382979e-05 5.673759e-05 3.829787e-05 [36] 3.971631e-05 3.687943e-05 1.985816e-05 2.411348e-05 5.673759e-05 [41] 3.829787e-05 1.702128e-05 1.985816e-05 2.411348e-05 1.702128e-05 [46] 1.843972e-05 1.560284e-05 1.418440e-05 5.673759e-06 1.560284e-05 [51] 8.510638e-06 4.255319e-06 4.255319e-06 9.929078e-06 1.418440e-06 [56] 5.673759e-06 1.134752e-05 4.255319e-06 2.836879e-06 1.418440e-06 [61] 4.255319e-06 2.836879e-06 0.000000e+00 2.836879e-06 4.255319e-06 [66] 4.255319e-06 5.673759e-06 0.000000e+00 1.418440e-06 0.000000e+00 [71] 0.000000e+00 2.836879e-06 1.418440e-06 0.000000e+00 5.673759e-06 [76] 0.000000e+00 0.000000e+00 0.000000e+00 1.418440e-06 0.000000e+00 [81] 1.418440e-06 0.000000e+00 1.418440e-06 1.418440e-06 0.000000e+00 [86] 0.000000e+00 0.000000e+00 0.000000e+00 1.418440e-06 0.000000e+00 [91] 0.000000e+00 0.000000e+00 2.836879e-06 0.000000e+00 1.418440e-06 [96] 0.000000e+00 0.000000e+00 0.000000e+00 1.418440e-06 0.000000e+00 [101] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [106] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [111] 0.000000e+00 0.000000e+00 1.418440e-06 0.000000e+00 0.000000e+00 [116] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [121] 0.000000e+00 0.000000e+00 1.418440e-06 0.000000e+00 0.000000e+00 [126] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [131] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [136] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [141] 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 0.000000e+00 [146] 0.000000e+00 0.000000e+00 1.418440e-06

\$mids

[1] [12] [23] [34] [45] [56] [67] [78] [89] [100] 9950 10050 10150 10250 10350 10450 10550 10650 10750 10850 10950 [111] 11050 11150 11250 11350 11450 11550 11650 11750 11850 11950 12050 [122] 12150 12250 12350 12450 12550 12650 12750 12850 12950 13050 13150 [133] 13250 13350 13450 13550 13650 13750 13850 13950 14050 14150 14250 [144] 14350 14450 14550 14650 14750

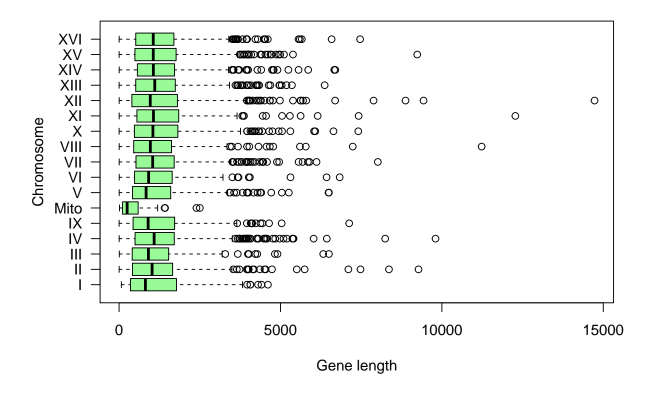


Figure 2: Boîte à moustache indiquant la distribution de longueur des gènes par chromosome.

Other types of graphs allow you to explore the distribution of a set of data. In particular, box plots display, for a series of data, the median, the quarterfinal range, a confidence interval and outliers.

7. Descriptive parameters

Calculate the parameters of central tendency (mean, median, mode) and dispersion (variance, standard deviation, inter-quarterly deviation)

- for the genes of chromosome III;
- for all yeast genes.

[1] 194 1

[1] "data.frame"

[1] "numeric"

length1 length2 length3 length4 length5 length6 741 1845 1374 780 630 525

[1] "Chromosome III contains 194 CDS"

[1] 1169.521

Ah ah! (skeptical tone) The R function sd() does not compute the standard deviation of the input numbers (s), but the estimate of the standard deviation of the population $(\hat{\sigma})$

Display these parameters on the histogram of gene length, using the function arrows()

8. Intervalle de confiance

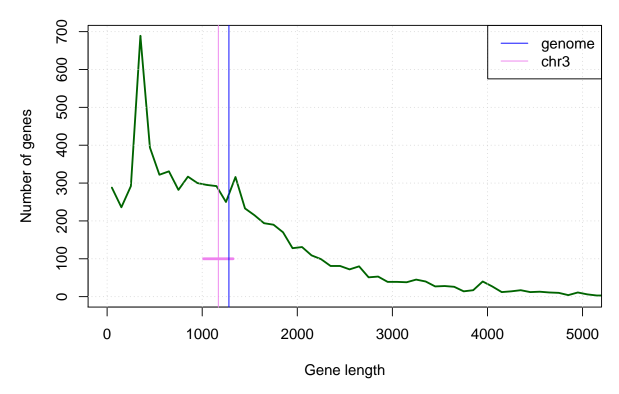
From genes of chromosome III (considered as the sample available in 1992), calculate a confidence interval around the mean, and formulate the interpretation of this confidence interval. Then evaluate whether or not this confidence interval covered the average population (all genes in the yeast genome, which became available 4 years after chromosome III).

$$\bar{x} \pm \frac{\hat{\sigma}}{\sqrt(n)} \cdot t_{1-\alpha/2}^{n-1}$$

[1] -1.972332

Draw a polygon of frequencies indicating the number of genes per class (class medium).

Frequency polygon



9. Distribution of gene length

- From the result of hist(), retrieve an array (in a variable of type data.frame) indicating the absolute frequencies (count) according to the median class size (mids),
- Add to this table a column indicating the relative frequency of each class of gene length.
- Add columns to this table indicating the **empirical distribution function** gene lengths (number of genes of a size less than or equal to each observed x value, and relative frequency of this number).
 - basic function: cumsum()
 - advanced function:ecdf()
- by using the functions plot() and lines(), draw a graph representing the absolute frequency per class (medians of classes in X, counts in Y), and the empirical distribution function.
 - suggestion: superposez les ??utilisez le type de lignes "h" pour les fréquences de classe, et "l" ou "s" pour la fonction de répartition.

10. Randomly expected distribution for gene length

Based on the genome size (12.156.679 bp) and codon genomic frequencies defined below, calculate the random expected gene length distribution, and add it to the graph.

You can download the genomic frequencies of all polynucleotides here: 3nt_genomic_Saccharomyces_cerevisiae-ovlp-1str.tab

Alternative: create a variable freq.3nt and manually assign the values for the 4? required polynucleotides from the table below.

sequence	frequency	occurrences
AAA	0.0394	478708
ATG	0.0183	221902
TAA	0.0224	272041
TAG	0.0129	156668
TGA	0.0201	244627

11. Before finishing: keep track of your session

Tractability is an essential issue in science. The function R sessionInfo() provides a summary of the conditions of a work session: version of R, operator system, libraries of functions used.

sessionInfo()

R version 3.6.1 (2019-07-05)

Platform: x86_64-apple-darwin15.6.0 (64-bit)

Running under: macOS Mojave 10.14.6

Matrix products: default

BLAS: /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib LAPACK: /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRlapack.dylib

locale:

[1] en US.UTF-8/en US.UTF-8/en US.UTF-8/c/en US.UTF-8/en US.UTF-8

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

[1] knitr_1.23

loaded via a namespace (and not attached):

- [1] compiler_3.6.1 magrittr_1.5 tools_3.6.1 htmltools_0.3.6 [5] yaml_2.2.0 Rcpp_1.0.2 stringi_1.4.3 rmarkdown_1.14 [9] highr_0.8 stringr_1.4.0 xfun_0.8 digest_0.6.20
- [13] evaluate_0.14