### Introducing ShortRead

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#### Plant genomes

- Arabidopsis thaliana is a small flowering plant
- First plant to have its genome sequenced
- Genome size 135 megabase pairs (Mbp)



#### Sequencing companies















<sup>&</sup>lt;sup>1</sup> Dan Koboldt massgenomics.org



#### fastq vs fasta

#### fastq

```
@ unique sequence identifier
```

raw sequence string

+ optional id

quality encoding per sequence letter

• fastq, fq

#### fasta

> unique sequence identifier

raw sequence string

• fasta, fa, seq

#### fasta

```
library(ShortRead)
# read fasta
fasample <- readFasta(dirPath = "data/", pattern = "fasta")</pre>
# print fasample
print(fasample)
class: ShortRead
length: 500 reads; width: 50 cycles
# methods accessors
methods(class = "ShortRead")
# Write a ShortRead object
writeFasta(fasample, file = "data/sample.fasta")
```

#### fastq

```
library(ShortRead)
# read fastq
fqsample <- readFastq(dirPath = "data/", pattern = "fastq")</pre>
# print fqsample
fqsample
class: ShortReadQ
length: 500 reads; width: 50 cycles
# methods accessors
methods(class = "ShortReadQ")
# Write a ShortRead object
writeFastq(fqsample, file = "data/sample.fastq.gz")
```

#### fastq sample

```
library(ShortRead)
# set the seed to draw the same read sequences every time
set.seed(123)
# Subsample of 500 bases
sampler <- FastqSampler("data/SRR1971253.fastq", 500)</pre>
# save the yield of 500 read sequences
sample_small <- yield(sampler)</pre>
# Class ShortReadQ
class(sample_small)
# length 500 reads
length(sample_small)
```

# You are ready!

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# Sequence quality

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#### Quality scores - Phred table

Quality value	Chance is wrong	Accuracy (%)
10	1 in 10	90
20	1 in 100	99
30	1 in 1 000	99.9
40	1 in 10 000	99.99
50	1 in 100 000	99.999

#### Encoding - Phred +33

```
# quality encoding
encoding(quality(fqsample))
```

#### Encoding characters and their scores

```
! " # $ % & ' ( ) * + , - . # encoding
0 1 2 3 4 5 6 7 8 9 10 11 12 13 # score

/ 0 1 2 3 4 5 6 7 8 9 : ; < # encoding
14 15 16 17 18 19 20 21 22 23 24 25 26 27 # score

= > ? @ A B C D E F G H I # encoding
28 29 30 31 32 33 34 35 36 37 38 39 40 # score
```

#### fastq quality

```
library(ShortRead)
quality(fqsample)
```

```
library(ShortRead)
sread(fqsample)[1]
# Quality is represented with ASCII characters
quality(fqsample)[1]
```

```
## PhredQuality instance
pq <- PhredQuality(quality(fqsample))
# transform encoding into scores
qs <- as(pq, "IntegerList")
qs # print scores</pre>
```

30 31 31 35 35 35 35 35 37 35 37 35 39 36 29 32 39 37 36 38 37 40 40 36 33 38 35 33 39 39 27 18 37 36 33 36 36 36 36 38 38 40 38 40 40 38 39 38 39 34



#### Quality assessment

```
library(ShortRead)
# Quality assessment
qaSummary <- qa(fqsample, lane = 1)  # optional lane
# class: ShortReadQQA(10)
# Names accessible with the quality assessment summary
names(qaSummary)</pre>
```

```
# Get a HTML report
browseURL(report(qaSummary))
```

```
library(ShortRead)
# sequences alphabet
alphabet(sread(fullSample))
```

```
A, C, G, T, M, R, W, S, Y, K, V, H, D, B, N, -, +, .
```

```
A C G T cycle
16839 16335 16740 10878 1
13056 13327 12064 22389 2
13666 15617 13198 18355 3
14723 15439 14239 16435 4
```



# Are you excited?

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#### Match and filter

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#### Duplicate sequences

- Biological sequence duplicates occur in nature
- Amplification from the steps in library preparation (PCR)
- Sequencing the sample more than once

Remove duplicates or at least mark them

Whole genome sequencing or exome sequencing

Mark duplicates using a threshold

RNA-seq and ChIP-seq

#### srduplicated

```
library(ShortRead)
# Counting duplicates TRUE is the number of duplicates
table(srduplicated(dfqsample))
FALSE TRUE
500
      500
# Cleaning reads from duplicates x[fun(x)]
cleanReads <- mydReads[srduplicated(mydReads) == FALSE]</pre>
# Counting duplicates
table(srduplicated(cleanReads))
FALSE
500
```



#### Creating your own filters

srFilter to filter based on a condition x[fun(x)]

Filter example

#### nFilter

```
library(ShortRead)
# save your filter, .name is optional
myFilter <- nFilter(threshold = 10, .name = "cleanNFilter")</pre>
# use the filter at reading point
filtered <- readFastq(dirPath = "data",
                      pattern = ".fastq",
                       filter = myFilter)
# you will retrieve only those reads that have a maximum of 10 N's
filtered
```

#### idFilter and polynFilter

```
library(ShortRead)
#id filter example
myFilterID <- idFilter(regex = ":3:1")</pre>
# will return only those ids that contain the regular expression
# optional parameters are .name, fixed and exclude
# use the filter at reading point
filtered <- readFastq(dirPath = "data", pattern = ".fastq",
                       filter = myFilterID)
# filter to remove poly-A regions
myFilterPolyA <- polynFilter(threshold = 10, nuc = c("A"))</pre>
# will return the sequences that have a maximun number of 10 consecutive A's
# use the filter for subsetting
filtered[myFilterPolyA(filtered)]
```

# Let's practice using filters!

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# Multiple and parallel sequence quality assessment

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#### Rqc

#### library(Rqc)

- Uses Bioconductor packages that you have already used:
  - o Biostrings, IRanges, methods, S4vectors
- New packages to discover in the following Bioconductor courses:
  - Rsamtools, GenomicAlignments, GenomicFiles, BiocParallel
- CRAN packages:
  - Knitr, dplyr, markdown, ggplot2, digest, shiny and Rccp

#### rqcQA

```
library(Rqc)
files <- # get the full path of the files you want to assess
qaRqc <- rqcQA(files)</pre>
# exploring qaRqc
class(qaRqc) # "list"
names(qaRqc) # name of the input files
# for each file
qaRqc[1]
# the class of the results is RqcResultSet
```

#### rqcQA arguments

```
library(Rqc)
# get the path of the files you want to assess
files <- "data/seq1.fq" "data/seq2.fq" "data/seq3.fq" "data/se4.fq"
qaRqc <- rqcQA(files, workers = 4))</pre>
# sample of sequences
set.seed(1111)
qaRqc_sample <- rqcQA(files, workers = 4, sample = TRUE, n = 500))</pre>
# paired-end files
pfiles <- "data/seq_11.fq" "data/seq1_2.fq" "data/seq2_1.fq" "data/seq2_2.fq"</pre>
qaRqc_paired <- rqcQA(pfiles, workers = 4, pair = c(1, 1, 2, 2)))
```

#### rqcReport and rqcResultSet

```
# create a report
reportFile <- rqcReport(qaRqc, templateFile = "myReport.Rmd")
browseURL(reportFile)
#The class of qaRqc is rqcResultSet
methods(class = "RqcResultSet")</pre>
```

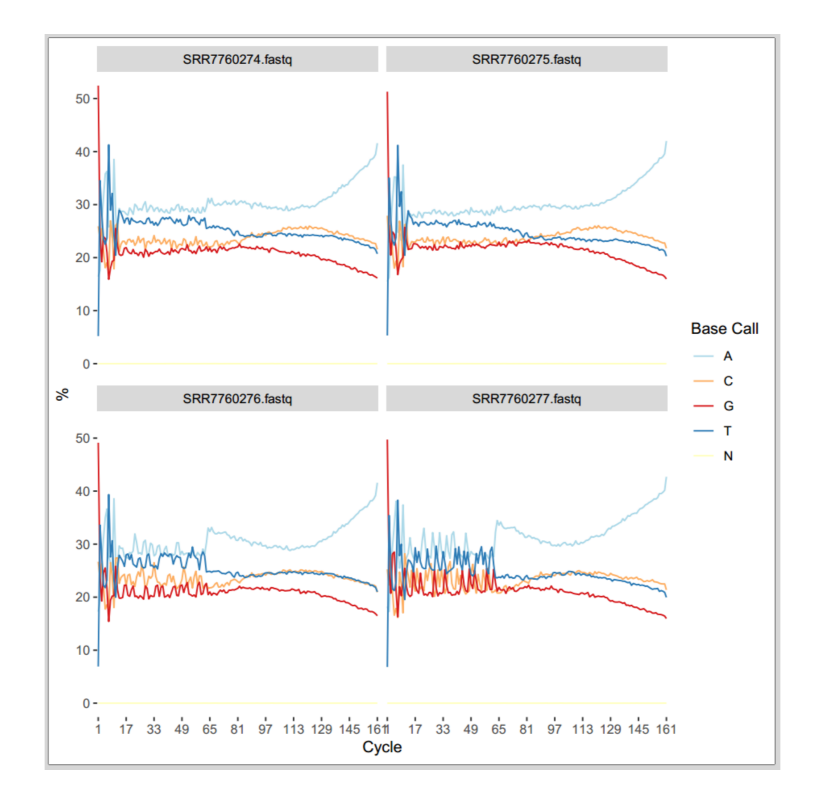
#### perFileInformation

```
qaRqc <- rqcQA(files, workers = 4))
perFileInformation(qaRqc)</pre>
```

```
filename
                            group reads total.reads
                pair format
                                                    path
                    FASTQ
                             None 1e+06
                                           2404795 ./data
SRR7760274.fastq 1
SRR7760275.fastq 2
                    FASTQ
                             None 1e+06
                                           1508139 ./data
                             None 1e+06
SRR7760276.fastq 3
                    FASTQ
                                           1950463 ./data
                                           2629588 ./data
SRR7760277.fastq 4
                    FASTQ
                             None 1e+06
```

#### Plot functions

rqc Plot functions	rqc Plot functions	
rqcCycleAverageQualityPcaPlot()	rqcGroupCycleAverageQualityPlot()	
rqcCycleAverageQualityPlot()	rqcReadQualityBoxPlot()	
rqcCycleBaseCallsLinePlot()	rqcReadQualityPlot()	
rqcCycleBaseCallsPlot()	rqcReadWidthPlot()	
rqcCycleGCPlot()	rqcReadFrequencyPlot()	
rqcCycleQualityBoxPlot()	rqcCycleQualityPlot()	





# You are ready!

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## Congratulations!

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#### You learned...

- Install packages from Bioconductor by using the BiocInstaller package.
- Techniques for reading, manipulating and filtering raw genomic data using BioStrings, GenomicRanges and ShortRead.
- To work with BSgenome and TxDb built-in datasets. Then used these to identify patterns by using matching functions.
- Check the quality of sequence files using ShortRead and Rqc.

#### You explored



# Keep learning!

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