# EDASeq: Exploratory Data Analysis and Normalization for RNA-Seq 

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## Modified: April 25, 2015. Compiled: October 30, 2017

## Contents

$\qquad$1 Introduction1
2 Reading in unaligned and aligned read data ..... 2
3 Read-level EDA. ..... 3
4 Gene-level EDA. ..... 5
5 Normalization ..... 9
6 Differential expression analysis ..... 10
6.1 edgeR ..... 11
6.2 DESeq ..... 11
7 Definitions and conventions ..... 12
7.1 Rounding ..... 12
7.2 Zero counts ..... 12
7.3 Offset ..... 13
8 Retrieving gene length and GC-content ..... 14
9 SessionInfo ..... 14

## 1 Introduction

In this document, we show how to conduct Exploratory Data Analysis (EDA) and normalization for a typical RNA-Seq experiment using the package EDASeq.

One can think of EDA for RNA-Seq as a two-step process: "read-level" EDA helps in discovering lanes with low sequencing depths, quality issues, and unusual nucleotide frequencies, while "gene-level" EDA can capture mislabeled lanes, issues with distributional assumptions (e.g., over-dispersion), and GC-content bias.

The package also implements both "within-lane" and "between-lane" normalization procedures, to account, respectively, for within-lane gene-specific (and possibly lane-specific) effects on read counts (e.g., related to gene length or GC-content) and for between-lane distributional differences in read counts (e.g., sequencing depths).
To illustrate the functionality of the EDASeq package, we make use of the Saccharomyces cerevisiae RNA-Seq data from Lee et al. [1]. Briefly, a wild-type strain and three mutant strains were sequenced using the Solexa 1G Genome Analyzer. For each strain, there are four technical replicate lanes from the same library preparation. The reads were aligned using Bowtie [2], with unique mapping and allowing up to two mismatches.

The leeBamViews package provides a subset of the aligned reads in BAM format. In particular, only the reads mapped between bases 800,000 and 900,000 of chromosome XIII are considered. We use these reads to illustrate read-level EDA.
The yeastRNASeq package contains gene-level read counts for four lanes: two replicates of the wild-type strain ("wt") and two replicates of one of the mutant strains ("mut"). We use these data to illustrate gene-level EDA.

```
require(EDASeq)
require(yeastRNASeq)
require(leeBamViews)
```


## 2 Reading in unaligned and aligned read data

Unaligned reads. Unaligned (unmapped) reads stored in FASTQ format may be managed via the class FastqFileList imported from ShortRead. Information related to the libraries sequenced in each lane can be stored in the elementMetadata slot of the FastqFileList object.

```
files <- list.files(file.path(system.file(package = "yeastRNASeq"),
    "reads"), pattern = "fastq", full.names = TRUE)
names(files) <- gsub("\\.fastq.*", "", basename(files))
met <- DataFrame(conditions=c(rep("mut",2), rep("wt",2)),
        row.names=names(files))
fastq <- FastqFileList(files)
elementMetadata(fastq) <- met
fastq
## FastqFileList of length 4
## names(4): mut_1_f mut_2_f wt_1_f wt_2_ff
```

Aligned reads. The package can deal with aligned (mapped) reads in BAM format, using the class BamFileList from Rsamtools. Again, the elementMetadata slot can be used to store lane-level sample information.

```
files <- list.files(file.path(system.file(package = "leeBamViews"), "bam"),
    pattern = "bam$", full.names = TRUE)
names(files) <- gsub("\\.bam", "", basename(files))
```

```
gt <- gsub(".*/", "", files)
gt <- gsub("__**", "", gt)
lane <- gsub(".*(.)$", "\\1", gt)
geno <- gsub(".$", "", gt)
pd <- DataFrame(geno=geno, lane=lane,
    row.names=paste(geno,lane,sep="."))
bfs <- BamFileList(files)
elementMetadata(bfs) <- pd
bfs
## BamFileList of length 8
## names(8): isowt5_13e isowt6_13e ... xrn1_13e xrn2_13e
```


## 3 Read-level EDA

Numbers of unaligned and aligned reads. One important check for quality control is to look at the total number of reads produced in each lane, the number and the percentage of reads mapped to a reference genome. A low total number of reads might be a symptom of low quality of the input RNA, while a low mapping percentage might indicate poor quality of the reads (low complexity), problems with the reference genome, or mislabeled lanes.

```
colors <- c(rep(rgb (1,0,0,alpha=0.7),2),
    rep (rgb (0,0,1,alpha=0.7),2),
    rep(rgb (0,1,0,alpha=0.7),2),
    rep(rgb (0,1,1,alpha=0.7),2))
barplot(bfs,las=2,col=colors)
plotQuality(bfs,col=colors,lty=1)
legend("topright",unique(elementMetadata(bfs)[,1]), fill=unique(colors))
```


(a) Number of mapped reads

Figure 1: Per-lane number of mapped reads and quality scores

Figure 1a, produced using the barplot method for the BamFileList class, displays the number of mapped reads for the subset of the yeast dataset included in the package leeBamViews. Unfortunately, leeBamViews does not provide unaligned reads, but barplots of the total number of reads can be obtained using the barplot method for the

FastqFileList class. Analogously, one can plot the percentage of mapped reads with the plot method with signature $c(x=$ "BamFileList", $y=$ "FastqFileList"). See the manual pages for details.

Read quality scores. As an additional quality check, one can plot the mean per-base (i.e., per-cycle) quality of the unmapped or mapped reads in every lane (Figure 1b).

Individual lane summaries. If one is interested in looking more thoroughly at one lane, it is possible to display the per-base distribution of quality scores for each lane (Figure 2a) and the number of mapped reads stratified by chromosome (Figure 2b) or strand. As expected, all the reads are mapped to chromosome XIII.
plotQuality (bfs[[1]], cex.axis=.8) barplot (bfs[[1]],las=2)


Figure 2: Quality scores and number of mapped reads for lane 'isowt5_13e.'

Read nucleotide distributions. A potential source of bias is related to the sequence composition of the reads. The function plotNtFrequency plots the per-base nucleotide frequencies for all the reads in a given lane (Figure 3).
plotNtFrequency (bfs[[1]])


Figure 3: Per-base nucleotide frequencies of mapped reads for lane 'isowt5_13e.'

## 4 Gene-level EDA

Examining statistics and quality metrics at a read level can help in discovering problematic libraries or systematic biases in one or more lanes. Nevertheless, some biases can be difficult to detect at this scale and gene-level EDA is equally important.

Classes and methods for gene-level counts. There are several Bioconductor packages for aggregating reads over genes (or other genomic regions, such as, transcripts and exons) given a particular genome annotation, e.g., IRanges, ShortRead, Genominator, Rsubread. See their respective vignettes for details.

Here, we consider this step done and load the object geneLevelData from yeastRNASeq, which provides gene-level counts for 2 wild-type and 2 mutant lanes from the yeast dataset of Lee et al. [1] (see the Genominator vignette for an example on the same dataset).

```
data(geneLevelData)
head(geneLevelData)
\begin{tabular}{lrrrr} 
\#\# & mut_1 & mut_2 & wt_1 & wt_2 \\
\#\# & YHR055C & 0 & 0 & 0 \\
\#\# YPR161C & 38 & 39 & 35 & 34 \\
\#\# YOL138C & 31 & 33 & 40 & 26 \\
\#\# YDR395W & 55 & 52 & 47 & 47 \\
\#\# YGR129W & 29 & 26 & 5 & 5 \\
\#\# YPR165W & 189 & 180 & 151 & 180
\end{tabular}
```

Since it is useful to explore biases related to length and GC-content, the EDASeq package provides, for illustration purposes, length and GC-content for S. cerevisiae genes (based on SGD annotation, version r64 [3]). Functionality for automated retrieval of gene length and GC-content is introduced in the last section of the vignette.

```
data(yeastGC)
head(yeastGC)
## YAL001C YAL002W YAL003W YAL004W YAL005C YAL007C
## 0.3712317 0.3717647 0.4460548 0.4490741 0.4406428 0.3703704
data(yeastLength)
head(yeastLength)
## YAL001C YAL002W YALO03W YALO04W YALO05C YALO07C
## 3483 3825 % 621 
```

First, we filter the non-expressed genes, i.e., we consider only the genes with an average read count greater than 10 across the four lanes and for which we have length and GCcontent information.

```
filter <- apply(geneLevelData,1,function(x) mean(x)>10)
table(filter)
## filter
## FALSE TRUE
## 1988 5077
```

```
common <- intersect(names(yeastGC),
    rownames(geneLevelData[filter,]))
length(common)
## [1] 4994
```

This leaves us with 4994 genes.
The EDASeq package provides the SeqExpressionSet class to store gene counts, (lane-level) information on the sequenced libraries, and (gene-level) feature information. We use the data frame met created in Section 2 for the lane-level data. As for the feature data, we use gene length and GC-content.

```
feature <- data.frame(gc=yeastGC,length=yeastLength)
data <- newSeqExpressionSet(counts=as.matrix(geneLevelData[common,]),
        featureData=feature[common,],
        phenoData=data.frame(
            conditions=c(rep("mut",2),rep("wt",2)),
            row.names=colnames(geneLevelData)))
data
## SeqExpressionSet (storageMode: lockedEnvironment)
## assayData: }4994\mathrm{ features, 4 samples
## element names: counts, normalizedCounts, offset
## protocolData: none
## phenoData
## sampleNames: mut_1 mut_2 wt_1 wt_2
## varLabels: conditions
## varMetadata: labelDescription
## featureData
## featureNames: YAL001C YAL002W ... YPR201W (4994
## total)
## fvarLabels: gc length
## fvarMetadata: labelDescription
## experimentData: use 'experimentData(object)'
## Annotation:
```

Note that the row names of counts and featureData must be the same; likewise for the row names of phenoData and the column names of counts. As in the CountDataSet class, the expression values can be accessed with counts, the lane information with pData, and the feature information with $£$ Data.

```
head(counts(data))
## mut_1 mut_2 wt_1 wt_2
## YAL001C 80 83 27 40
## YALO02W 33 38 53 66
## YALO03W 1887 1912 270 270
## YALO04W 90 110 276 295
## YALOO5C 325 316 874 935
## YALOO7C 27 30 19 24
pData(data)
```

```
## conditions
## mut_1 mut
## mut_2 mut
## wt_1 wt
## wt_2 wt
head(fData(data))
\begin{tabular}{lrr} 
\#\# & gc & length \\
\#\# YAL001C & 0.3712317 & 3483 \\
\#\# YAL002W & 0.3717647 & 3825 \\
\#\# YAL003W & 0.4460548 & 621 \\
\#\# YAL004W & 0.4490741 & 648 \\
\#\# YAL005C & 0.4406428 & 1929 \\
\#\# YAL007C & 0.3703704 & 648
\end{tabular}
```

The SeqExpressionSet class has two additional slots: normalizedCounts and off set (matrices of the same dimension as counts), which may be used to store a matrix of normalized counts and of normalization offsets, respectively, to be used for subsequent analyses (see Section 5 and the edgeR vignette for details on the role of offsets). If not specified, the offset is initialized as a matrix of zeros.
head (offst (data))

| \#\# | mut_1 | mut_2 | wt_1 | wt_2 |
| :--- | ---: | ---: | ---: | ---: |
| \#\# YAL001C | 0 | 0 | 0 | 0 |
| \#\# YAL002W | 0 | 0 | 0 | 0 |
| \#\# YAL003W | 0 | 0 | 0 | 0 |
| \#\# YAL004W | 0 | 0 | 0 | 0 |
| \#\# YAL005C | 0 | 0 | 0 | 0 |
| \#\# YAL007C | 0 | 0 | 0 | 0 |

Between-lane distribution of gene-level counts. One of the main considerations when dealing with gene-level counts is the difference in count distributions between lanes. The boxplot method provides an easy way to produce boxplots of the logarithms of the gene counts in each lane (Figure 4).
boxplot (data, col=colors[1:4])


Figure 4: Between-lane distribution of gene-level counts (log)
The MDP lot method produces a mean-difference plot (MD-plot) of read counts for two lanes (Figure 5).

MDPlot (data, c(1, 3))


Figure 5: Mean-difference plot of the gene-level counts (log) of lanes 'mut_1' and 'wt_1.'

Over-dispersion. Although the Poisson distribution is a natural and simple way to model count data, it has the limitation of assuming equality of the mean and variance. For this reason, the negative binomial distribution has been proposed as an alternative when the data show over-dispersion. The function meanVarPlot can be used to check whether the count data are over-dispersed (for the Poisson distribution, one would expect the points in Figures 6 to be evenly scattered around the black line).
meanVarPlot (data[,1:2], log=TRUE, ylim=c (0,16))
meanVarPlot(data, log=TRUE, $\operatorname{ylim=c}(0,16)$ )


Figure 6: Mean-variance relationship for the two mutant lanes and all four lanes: the black line corresponds to the Poisson distribution (variance equal to the mean), while the red curve is a lowess fit

Note that the mean-variance relationship should be examined within replicate lanes only (i.e., conditional on variables expected to contribute to differential expression). For the yeast dataset, it is not surprising to see no evidence of over-dispersion for the two mutant technical replicate lanes (Figure 6a); likewise for the two wild-type lanes. However, one expects over-dispersion in the presence of biological variability, as seen in Figure 6b when considering at once all four mutant and wild-type lanes [4-6].

Gene-specific effects on read counts. Several authors have reported selection biases related to sequence features such as gene length, GC-content, and mappability [4, 7-9].

In Figure 7, obtained using biasplot, one can see the dependence of gene-level counts on GC-content. The same plot could be created for gene length or mappability instead of GC-content.
biasPlot(data, "gc", log=TRUE, ylim=c(1,5))


Figure 7: Lowess regression of the gene-level counts (log) on GC-content for each lane, color-coded by experimental condition

To show that GC-content dependence can bias differential expression analysis, one can produce stratified boxplots of the log-fold-change of read counts from two lanes using the biasBoxplot method (Figure 8). Again, the same type of plots can be created for gene length or mappability.
$\operatorname{lfc}<-\log (\operatorname{counts}($ data $[, 3]+0.1)-\log (\operatorname{counts}($ data) $[, 1]+0.1)$
biasBoxplot(lfc, fData(data) \$gc)


Figure 8: Boxplots of the log-fold-change between 'mut_1' and 'wt_1' lanes stratified by GC-content

## 5 Normalization

Following Risso et al. [8], we consider two main types of effects on gene-level counts: (1) within-lane gene-specific (and possibly lane-specific) effects, e.g., related to gene length or GC-content, and (2) effects related to between-lane distributional differences, e.g., sequencing depth. Accordingly, withinLaneNormalization and betweenLa neNormalization adjust for the first and second type of effects, respectively. We recommend to normalize for within-lane effects prior to between-lane normalization.
We implemented four within-lane normalization methods, namely: loess robust local regression of read counts (log) on a gene feature such as GC-content (loess), globalscaling between feature strata using the median (median), global-scaling between feature strata using the upper-quartile (upper), and full-quantile normalization between feature strata (full). For a discussion of these methods in context of GC-content normalization see Risso et al. [8].

```
dataWithin <- withinLaneNormalization(data,"gc", which="full")
dataNorm <- betweenLaneNormalization(dataWithin, which="full")
```

Regarding between-lane normalization, the package implements three of the methods introduced in Bullard et al. [4]: global-scaling using the median (median), global-scaling using the upper-quartile (upper), and full-quantile normalization (full).
Figure 9 shows how after full-quantile within- and between-lane normalization, the GCcontent bias is reduced and the distribution of the counts is the same in each lane.

```
biasPlot(dataNorm, "gc", log=TRUE, ylim=c(1,5))
```

boxplot(dataNorm, col=colors)


Figure 9: Full-quantile within- and between-lane normalization
(a) Lowess regression of normalized gene-level counts (log) on GC-content for each lane. (b) Between-lane distribution of normalized gene-level counts (log).

Offset. Some authors have argued that it is better to leave the count data unchanged to preserve their sampling properties and instead use an offset for normalization purposes in the statistical model for read counts [5, 6, 9]. This can be achieved easily using the argument offset in both normalization functions.

```
dataOffset <- withinLaneNormalization(data,"gc",
    which="full",offset=TRUE)
dataOffset <- betweenLaneNormalization(dataOffset,
    which="full",offset=TRUE)
```

Note that the dataOffset object will have both normalized counts and offset stored in their respective slots.

## 6 Differential expression analysis

One of the main applications of RNA-Seq is differential expression analysis. The normalized counts (or the original counts and the offset) obtained using the EDASeq package can be supplied to packages such as edgeR [5] or DESeq [6] to find differentially expressed genes. This section should be considered only as an illustration of the compatibility of the results of EDASeq with two of the most widely used packages for differential expression; our aim is not to compare differential expression strategies (e.g., normalized counts vs. offset).

## 6.1 edgeR

We can perform a differential expression analysis with edgeR based on the original counts by passing an offset to the generalized linear model. For simplicity, we estimate a common dispersion parameter for all genes. See the edge $R$ vignette for details about how to perform a differential expression analysis using a gene-specific dispersion or more complex designs.

```
library(edgeR)
design <- model.matrix(~conditions, data=pData(dataOffset))
disp <- estimateGLMCommonDisp(counts(dataOffset),
    design, offset=-offst(dataOffset))
fit <- glmFit(counts(dataOffset), design, disp, offset=-offst(dataOffset))
lrt <- glmLRT(fit, coef=2)
topTags(lrt)
## Coefficient: conditionswt
## logFC logCPM LR PValue
## YGL088W - 5.553632 27.34855 924.5709 4.471753e-203
## YPL198W - 7.425857 26.51941 899.9077 1.027736e-197
## YMR013W-A -6.236486 26.16157 775.8177 9.768538e-171
## YHR156C -8.132765 25.10950 647.7490 6.896873e-143
## YLR106C -4.162281 25.93311 613.0509 2.427567e-135
## YGL076C -4.206669 26.10029 585.7168 2.140773e-129
## YOR040W -4.974040 25.46010 567.7222 1.757105e-125
## YBLO04W -5.261038 25.03227 556.3022 5.358092e-123
## YALO03W -2.991170 27.49021 517.8531 1.240623e-114
## YJLO47C-A -4.346547 24.96897 416.9532 1.123627e-92
## FDR
## YGL088W 2.233193e-199
## YPL198W 2.566256e-194
## YMR013W-A 1.626136e-167
## YHR156C 8.610746e-140
## YLR106C 2.424653e-132
## YGL076C 1.781837e-126
## YOR040W 1.253569e-122
## YBLO04W 3.344789e-120
## YALO03W 6.884082e-112
## YJL047C-A 5.611395e-90
```


### 6.2 DESeq

We can perform a differential expression analysis with DESeq based on the normalized counts by using the coerce method from the SeqExpressionSet class to the CountDataSet class of DESeq. When working with data that have been normalized for both within- and between-lane effects, we force the size factors to be one, since differences
in lane sequencing depths have already been accounted for in our between-lane normalization. One could also consider only within-lane normalization and account for differences in sequencing depth by estimating the size factors using $D E S e q$.

```
library(DESeq)
counts <- as(dataNorm, "CountDataSet")
sizeFactors(counts) <- rep(1,4)
counts <- estimateDispersions(counts)
res <- nbinomTest(counts, "wt", "mut")
head(res)
## id baseMean baseMeanA baseMeanB foldChange
## 1 YAL001C 92.00 58.0 126.0 2.1724138
## 2 YALO02W 73.75 111.0 36.5 0.3288288
## 3 YALO03W 565.00 126.0 1004.0 7.9682540
## 4 YALO04W 100.00 133.0 67.0 0.5037594
## 5 YALO05C 259.25 330.5 188.0 0.5688351
## 6 YAL007C 32.25 37.0 27.5 0.7432432
## log2FoldChange pval padj
## 1 1.1192989 3.415271e-04 3.732136e-03
## 2 -1.6045913 2.106666e-06 4.826006e-05
## 3 2.9942636 3.882861e-31 1.615917e-28
## 4 -0.9891932 1.181444e-03 1.022553e-02
## 5 -0.8139176 1.418425e-03 1.178638e-02
## 6 -0.4280937 3.346353e-01 5.776740e-01
```


## 7 Definitions and conventions

### 7.1 Rounding

After either within-lane or between-lane normalization, the expression values are not counts anymore. However, their distribution still shows some typical features of counts distribution (e.g., the variance depends on the mean). Hence, for most applications, it is useful to round the normalized values to recover count-like values, which we refer to as "pseudo-counts".

By default, both withinLaneNormalization and betweenLaneNormalization round the normalized values to the closest integer. This behavior can be changed by specifying round=FALSE. This gives the user more flexibility and assures that rounding approximations do not affect subsequent computations (e.g., recovering the offset from the normalized counts).

### 7.2 Zero counts

To avoid problems in the computation of logarithms (e.g. in log-fold-changes), we add a small positive constant (namely 0.1) to the counts. For instance, the log-fold-change between $y_{1}$ and $y_{2}$ is defined as

$$
\frac{\log \left(y_{1}+0.1\right)}{\log \left(y_{2}+0.1\right)}
$$

### 7.3 Offset

We define an offset in the normalization as

$$
o=\log \left(y_{\text {norm }}+0.1\right)-\log \left(y_{\text {raw }}+0.1\right),
$$

where $y_{\text {norm }}$ and $y_{\text {raw }}$ are the normalized and raw counts, respectively.
One can easily recover the normalized data from the raw counts and offset, as shown here:

```
dataNorm <- betweenLaneNormalization(data, round=FALSE, offset=TRUE)
norm1 <- normCounts(dataNorm)
norm2 <- exp(log(counts(dataNorm) + 0.1 ) + offst(dataNorm)) - 0.1
head(norm1 - norm2)
## mut_1 mut_2 wt_1
## YAL001C -1.421085e-14 -1.421085e-14 0.000000e+00
## YAL002W 3.552714e-15 -7.105427e-15 -2.131628e-14
## YAL003W 4.547474e-13 -2.273737e-13 1.136868e-13
## YALO04W -2.842171e-14 -1.421085e-14 5.684342e-14
## YALO05C -1.136868e-13 5.684342e-14 -3.410605e-13
## YAL007C 3.552714e-15 3.552714e-15 -3.552714e-15
## wt_2
## YAL001C 0.000000e+00
## YAL002W 0.000000e+00
## YAL003W -5.684342e-14
## YALO04W -5.684342e-14
## YALO05C -1.136868e-13
## YALO07C 3.552714e-15
```

Note that the small constant added in the definition of offset does not matter when pseudo-counts are considered, i.e.,

```
head(round(normCounts(dataNorm)) - round(counts(dataNorm) * exp(offst(dataNorm))))
```

| \#\# | mut_1 | mut_2 | wt_1 | wt_2 |
| :--- | ---: | ---: | ---: | ---: | ---: |
| \#\# YAL001C | 0 | 0 | 0 | 0 |
| \#\# YAL002W | 0 | 0 | 0 | 0 |
| \#\# YAL003W | 0 | 0 | 0 | 0 |
| \#\# YAL004W | 0 | 0 | 0 | 0 |
| \#\# YAL005C | 0 | 0 | 0 | 0 |
| \#\# YAL007C | 0 | 0 | 0 | 0 |

We defined the offset as the log-ratio between normalized and raw counts. However, the edgeR functions expect as offset argument the log-ratio between raw and normalized counts. One must use -offst (offsetData) as the offset argument of edgeR (see Section 6.1).

## 8 Retrieving gene length and GC-content

Two essential features the gene-level EDA normalizes for are gene length and GCcontent. As users might wish to automatically retrieve this information, we provide the function getGeneLengthAndGCContent. Given selected ENTREZ or ENSEMBL gene IDs and the organism under investigation, this can be done either based on BioMart (default) or using BioC annotation utilities.

```
getGeneLengthAndGCContent (id=c("ENSG00000012048", "ENSG00000139618"), org="hsa")
## length gc
## ENSG00000012048 8922 0.4588657
## ENSG00000139618 12273 0.3644586
```

Accordingly, we can retrieve the precalculated yeast data that has been used throughout the vignette via

```
fData(data) <- getGeneLengthAndGCContent(featureNames(data),
    org="sacCer3", mode="org.db")
```


## 9 Sessionlnfo

toLatex(sessionInfo())

- R version 3.4.2 (2017-09-28), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Running under: Ubuntu 16.04.3 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.6-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.6-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: BSgenome 1.46.0, Biobase 2.38.0, BiocGenerics 0.24.0, BiocParallel 1.12.0, Biostrings 2.46.0, DESeq 1.30.0, DelayedArray 0.4.0, EDASeq 2.12.0, GenomelnfoDb 1.14.0, GenomicAlignments 1.14.0, GenomicRanges 1.30.0, IRanges 2.12.0, Rsamtools 1.30.0, S4Vectors 0.16.0, ShortRead 1.36.0, SummarizedExperiment 1.8.0, XVector 0.18.0, edgeR 3.20.0, knitr 1.17, lattice 0.20-35, leeBamViews 1.13.1, limma 3.34.0, locfit 1.5-9.1, matrixStats 0.52.2, rtracklayer 1.38.0, yeastRNASeq 0.15.0
- Loaded via a namespace (and not attached): AnnotationDbi 1.40.0, BiocStyle 2.6.0, DBI 0.7, GenomeInfoDbData 0.99.1, GenomicFeatures 1.30.0, KernSmooth 2.23-15, Matrix 1.2-11, R.methodsS3 1.7.1, R.oo 1.21.0,
R.utils 2.5.0, R6 2.2.2, RColorBrewer 1.1-2, RCurl 1.95-4.8, RMySQL 0.10.13, RSQLite 2.0, Rcpp 0.12.13, XML 3.98-1.9, annotate 1.56.0, aroma.light 3.8.0, asserthat 0.2.0, backports 1.1.1, biomaRt 2.34.0, bit 1.1-12, bit64 0.9-7, bitops $1.0-6$, blob 1.1.0, compiler 3.4.2, digest 0.6 .12 , evaluate 0.10 .1 , genefilter 1.60.0, geneplotter 1.56.0, grid 3.4.2, highr 0.6 , htmltools 0.3.6, hwriter 1.3.2, latticeExtra $0.6-28$, magrittr 1.5 , memoise 1.1.0, prettyunits 1.0 .2 , progress 1.1.2, rlang 0.1 .2 , rmarkdown 1.6 , rprojroot 1.2 , splines 3.4 .2 , stringi 1.1.5, stringr 1.2.0, survival 2.41-3, tibble 1.3.4, tools 3.4.2, xtable 1.8-2, yaml 2.1.14, zlibbioc 1.24.0


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