

Manual for the package samExploreR

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Contents

1	Citation.	2
2	Dependencies.	2
2.1	For developers only.	2
3	Installation.	3
4	Quick start.	3
5	The samExplore function	5
6	exploreRob	11
7	exploreRep	11
8	plotSamExplorer	12
8.1	Plotting data using the function plotSamExplorer	13

1 Citation

If you use the samExploreR package, please cite [1] and [2].

2 Dependencies

SamExploreR depends on some R package available in Bioconductor that need to be installed before you can use the package. The symbol '>' indicates the R prompt.

```
> if (!requireNamespace("BiocManager", quietly=TRUE))
+   install.packages("BiocManager")
> BiocManager::install("samExploreR")
```

Installation command: call the following command from an R prompt.

2.1 For developers only

If a user wants to make changes to the package and rebuild it again, then the following packages need to be installed additionally. These are necessary for the compilation and the building the package.

```
> BiocManager::install(c("BiocCheck", "BiocGenerics", "RUnit"))
```

3 Installation

The samExploreR package is available from Bioconductor. To install the package, run the following commands:

Remark: To be added after uploaded to Bioconductor.

If you install it from a local download, run the following command in a terminal from the same directory where you downloaded the package:

4 Quick start

In the following, we demonstrate briefly the usage of the samExploreR package and the analysis procedures it provides. These examples require a BAM or SAM file from some RNA-seq experiment containing aligned reads. For the following examples, we provide the all necessary files, namely:

- aligned reads: Test.sam
- annotation file: Annot.gtf

In order to simulate 5 repeats (N_{boot}) for a virtual sequencing experiment with sequencing depth $f = 0.7$ execute:

```
> library(samExploreR)
> ## perform subsampling
> inpf <- RNAseqData.HNRNPC.bam.chr14_BAMFILES
> x <- samExplore(files=inpf,annot.inbuilt="hg19", subsample_d = 0.8)
```

NCBI RefSeq annotation for hg19 (build 37.2) is used.

```
=====
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===== ____ ) | |_) | |_) | | \ \ | |__ | ____ \ | | | |
===== |____/ \___/ |____/ | | \ \_____/ / \ \_____/
Rsubread 2.3.0
```

```
//===== featureCounts setting =====\
||                                                                 ||
```

Manual for the package samExploreR

```
||           Input files : 8 BAM files           ||
||           o ERR127306_chr14.bam             ||
||           o ERR127307_chr14.bam             ||
||           o ERR127308_chr14.bam             ||
||           o ERR127309_chr14.bam             ||
||           o ERR127302_chr14.bam             ||
||           o ERR127303_chr14.bam             ||
||           o ERR127304_chr14.bam             ||
||           o ERR127305_chr14.bam             ||
||
||           Annotation : inbuilt (hg19)        ||
||           Dir for temp files : .             ||
||           Threads : 1                       ||
||           Level : meta-feature level         ||
||           Paired-end : no                   ||
||           Multimapping reads : counted       ||
||           Multi-overlapping reads : not counted ||
||           Min overlapping bases : 1          ||
||
||=====||
||===== Running =====||
||
|| Load annotation file hg19_RefSeq_exon.txt ... ||
||           Features : 225074                 ||
||           Meta-features : 25702            ||
||           Chromosomes/contigs : 52         ||
||
|| Process BAM file ERR127306_chr14.bam...     ||
||           WARNING: Paired-end reads were found. ||
||           Total alignments : 800484         ||
||           Successfully assigned alignments : 680516 (85.0%) ||
||           Running time : 0.04 minutes       ||
||
|| Process BAM file ERR127307_chr14.bam...     ||
||           WARNING: Paired-end reads were found. ||
||           Total alignments : 875302         ||
||           Successfully assigned alignments : 744592 (85.1%) ||
||           Running time : 0.04 minutes       ||
||
|| Process BAM file ERR127308_chr14.bam...     ||
||           WARNING: Paired-end reads were found. ||
||           Total alignments : 875608         ||
||           Successfully assigned alignments : 741944 (84.7%) ||
||           Running time : 0.04 minutes       ||
```

```
||
|| Process BAM file ERR127309_chr14.bam...
||   WARNING: Paired-end reads were found.
||   Total alignments : 778464
||   Successfully assigned alignments : 661631 (85.0%)
||   Running time : 0.04 minutes
||
|| Process BAM file ERR127302_chr14.bam...
||   WARNING: Paired-end reads were found.
||   Total alignments : 742754
||   Successfully assigned alignments : 626250 (84.3%)
||   Running time : 0.04 minutes
||
|| Process BAM file ERR127303_chr14.bam...
||   WARNING: Paired-end reads were found.
||   Total alignments : 786196
||   Successfully assigned alignments : 659952 (83.9%)
||   Running time : 0.04 minutes
||
|| Process BAM file ERR127304_chr14.bam...
||   WARNING: Paired-end reads were found.
||   Total alignments : 810724
||   Successfully assigned alignments : 660986 (81.5%)
||   Running time : 0.04 minutes
||
|| Process BAM file ERR127305_chr14.bam...
||   WARNING: Paired-end reads were found.
||   Total alignments : 798376
||   Successfully assigned alignments : 658094 (82.4%)
||   Running time : 0.04 minutes
||
|| Write the final count table.
|| Write the read assignment summary.
||
||=====//
```

This results in a 5-dimensional list, whereas each component corresponds to a sub-sampled count vectors.

5 The samExplore function

The function `samExplore`, which is our modification of the `featureCounts` function of [2], performs a summerization of reads to genomic features of annotation.

Manual for the package samExploreR

The procedure of read subsampling works as follows: One of the input parameters is f , which is a fraction of reads that will be subsampled from the original SAM or BAM file,

$$f = \frac{\text{\#subsampled reads}}{\text{\#total reads}}. \quad \mathbf{1}$$

During the counts computing process every read, or a pair of reads for paired-end sequencing, is taken into account with probability f . This results in a reduction of the amount of reads and, therefore, the overall expression of the genes.

Further input arguments of the function are:

- files: names of SAM/BAM files to process
- annot_ext: annotation file
- isGTFAnnotationFile:
- GTF.featureType:
- N_boot: number of repeated experiments

As an output the function produces a vector of objects - results of featureCounts running.

```
> # Loading library
> library(samExploreR)
> # Performing subsampling
>
> inpf <- RNAseqData.HNRNPC.bam.chr14_BAMFILES
> # Performing robustness analysis for f = 0.7, number of replicates 5,
> #annotation entries 'gene', non-paired reads
> x <- samExplore(files=inpf,annot.inbuilt="hg19",GTF.featureType="exon",
+ GTF.attrType="gene_id", subsample_d = 0.8, N_boot=5)
```

NCBI RefSeq annotation for hg19 (build 37.2) is used.

```
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===== ____ ) | |__ | | |-) | | \ \ | |__ / ____ \ | | | |
===== |____/ \____/ |____/ | | \ \_____/ - / \ \_____/
Rsubread 2.3.0
```

```
//===== featureCounts setting =====\\
||
||          Input files : 8 BAM files          ||
||          o ERR127306_chr14.bam             ||
```

Manual for the package samExploreR

```
|| o ERR127307_chr14.bam ||
|| o ERR127308_chr14.bam ||
|| o ERR127309_chr14.bam ||
|| o ERR127302_chr14.bam ||
|| o ERR127303_chr14.bam ||
|| o ERR127304_chr14.bam ||
|| o ERR127305_chr14.bam ||
||
|| Annotation : inbuilt (hg19) ||
|| Dir for temp files : . ||
|| Threads : 1 ||
|| Level : meta-feature level ||
|| Paired-end : no ||
|| Multimapping reads : counted ||
|| Multi-overlapping reads : not counted ||
|| Min overlapping bases : 1 ||
||
||=====||
||===== Running =====||
||
|| Load annotation file hg19_RefSeq_exon.txt ... ||
|| Features : 225074 ||
|| Meta-features : 25702 ||
|| Chromosomes/contigs : 52 ||
||
|| Process BAM file ERR127306_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 800484 ||
|| Successfully assigned alignments : 680516 (85.0%) ||
|| Running time : 0.03 minutes ||
||
|| Process BAM file ERR127307_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 875302 ||
|| Successfully assigned alignments : 744592 (85.1%) ||
|| Running time : 0.04 minutes ||
||
|| Process BAM file ERR127308_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 875608 ||
|| Successfully assigned alignments : 741944 (84.7%) ||
|| Running time : 0.04 minutes ||
||
|| Process BAM file ERR127309_chr14.bam... ||
```

Manual for the package samExploreR

```

|| WARNING: Paired-end reads were found. ||
|| Total alignments : 778464 ||
|| Successfully assigned alignments : 661631 (85.0%) ||
|| Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127302_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 742754 ||
|| Successfully assigned alignments : 626250 (84.3%) ||
|| Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127303_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 786196 ||
|| Successfully assigned alignments : 659952 (83.9%) ||
|| Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127304_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 810724 ||
|| Successfully assigned alignments : 660986 (81.5%) ||
|| Running time : 0.04 minutes ||
|| ||
|| Process BAM file ERR127305_chr14.bam... ||
|| WARNING: Paired-end reads were found. ||
|| Total alignments : 798376 ||
|| Successfully assigned alignments : 658094 (82.4%) ||
|| Running time : 0.04 minutes ||
|| ||
|| Write the final count table. ||
|| Write the read assignment summary. ||
|| ||
\\=====//

```

```

> # Performing robustness analysis for f = 0.1, number of replicates 10,
> #annotation entries 'exon', paired reads
> x <- samExplore(files=inpfile,annot.inbuilt="hg19",GTF.featureType="gene",
+ GTF.attrType="gene_id", subsample_d = 0.8, N_boot=5)

```

NCBI RefSeq annotation for hg19 (build 37.2) is used.

```

=====
===== / ----| | | | - \ | -- \ | ----| / \ | -- \
===== | (--- | | | | | -) | |---) | |--- / \ | | | |
===== \--- \ | | | | - <| - / | -- | / \ \ | | | |
===== ----) | |---| | | -) | | \ \ | |---- / ---- \ | |--- |

```


Manual for the package samExploreR

```
===== |_____/ \_____/|_____/|_ | \_\_____/ / \_\_____/
Rsubread 2.3.0
```

```
//===== featureCounts setting =====\\
||
||      Input files : 8 BAM files
||          o ERR127306_chr14.bam
||          o ERR127307_chr14.bam
||          o ERR127308_chr14.bam
||          o ERR127309_chr14.bam
||          o ERR127302_chr14.bam
||          o ERR127303_chr14.bam
||          o ERR127304_chr14.bam
||          o ERR127305_chr14.bam
||
||      Annotation : inbuilt (hg19)
||      Dir for temp files : .
||      Threads : 1
||      Level : meta-feature level
||      Paired-end : no
||      Multimapping reads : counted
|| Multi-overlapping reads : not counted
||      Min overlapping bases : 1
||
||=====\\
```

```
//===== Running =====\\
||
|| Load annotation file hg19_RefSeq_exon.txt ...
||      Features : 225074
||      Meta-features : 25702
||      Chromosomes/contigs : 52
||
|| Process BAM file ERR127306_chr14.bam...
||      WARNING: Paired-end reads were found.
||      Total alignments : 800484
||      Successfully assigned alignments : 680516 (85.0%)
||      Running time : 0.03 minutes
||
|| Process BAM file ERR127307_chr14.bam...
||      WARNING: Paired-end reads were found.
||      Total alignments : 875302
||      Successfully assigned alignments : 744592 (85.1%)
||      Running time : 0.03 minutes
||
```

Manual for the package samExploreR

```
|| Process BAM file ERR127308_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 875608 ||
||   Successfully assigned alignments : 741944 (84.7%) ||
||   Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127309_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 778464 ||
||   Successfully assigned alignments : 661631 (85.0%) ||
||   Running time : 0.02 minutes ||
|| ||
|| Process BAM file ERR127302_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 742754 ||
||   Successfully assigned alignments : 626250 (84.3%) ||
||   Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127303_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 786196 ||
||   Successfully assigned alignments : 659952 (83.9%) ||
||   Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127304_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 810724 ||
||   Successfully assigned alignments : 660986 (81.5%) ||
||   Running time : 0.03 minutes ||
|| ||
|| Process BAM file ERR127305_chr14.bam... ||
||   WARNING: Paired-end reads were found. ||
||   Total alignments : 798376 ||
||   Successfully assigned alignments : 658094 (82.4%) ||
||   Running time : 0.03 minutes ||
|| ||
|| Write the final count table. ||
|| Write the read assignment summary. ||
|| ||
\\=====//
```

6 exploreRob

A cornerstone of any scientific study is the question regarding the reproducibility and robustness of obtained results. The function `exploreRob` allows the exploration of the robustness of results. It runs a standard one-way ANOVA test for groups of replicates, corresponding to different f values, for one fixed annotation. In this way, one can measure if the result of the analysis changes significantly with a change in the parameter sequencing depth, f .

The function `exploreRob` takes as input argument a data frame with the format shown in Table below. ??.

```
> data(df_sole)
> head(df_sole)
```

	Label	Variable	Value
1	New, Gene	0.05	3
2	New, Exon	0.05	1
3	Old, Gene	0.05	1
4	New, Gene	0.05	1
5	New, Exon	0.05	2
6	Old, Gene	0.05	0

In this table, the first column provides the labels for the annotation (`lbl`) used for the analysis, the second column gives the f value and the third column provides the value of the metric to be explored, e.g., the number of differentially expressed genes.

The function `exploreRob` splits this data frame up by considering only entries for one specific type of annotation. Then an ANOVA test is run for the groups of replicates that correspond to a given list of f values, see Fig. 1.

For instance, to explore the robustness of the annotation type `AnnotB` cross the f values 0.8, 0.9, 0.95 for the data frame `df` run:

```
> #Loading library
> library(samExploreR)
> data("df_sole")
> #Performing robustness analysis
> exploreRob(df_sole, lbl = 'New, Gene', f_vect = c(0.85, 0.9, 0.95))
```

7 exploreRep

Similar to `exploreRob`, the function `exploreRep` allows the exploration of the reproducibility of results. This function runs a one-way ANOVA test for groups of replicates, corresponding to one f value, across various annotation types. Thus, the influence of the annotation or the summarisation method can be explored.

Manual for the package samExploreR

`exploreRep` takes as input a data frame of form shown in Tab. 1. Here the first

	Label	Variable	Value
.	.	.	.
.	.	.	.
1	New, Gene	0.05	3
2	New, Exon	0.05	1
3	Old, Gene	0.05	1
4	New, Gene	0.05	1
5	New, Exon	0.05	2
6	Old, Gene	0.05	0
7	New, Gene	0.05	2
8	New, Exon	0.05	3
9	Old, Gene	0.05	0
10	New, Gene	0.05	2
.	.	.	.
.	.	.	.

Table 1: Input argument for the function `exploreRep`

column provides the labels for the annotation used for the analysis, the second column gives the f value and the third column provides the value of the metric to be explored, e.g., the number of differentially expressed genes.

`exploreRep` splits this data frame up to consider only results for one f value. ANOVA test is run for groups of replicates with corresponding to given list of annotation labels, Fig 1.

For instance, to explore the reproducibility for a f value of 0.7 cross the annotation types AnnotA, AnnotB, AnnotC for data frame `df` run:

```
> #Loading library
> library(samExploreR)
> data("df_0.9")
> #Performing robustness analysis
> t = exploreRep(df_0.9, lbl_vect = c('New, Gene', 'Old, Gene', 'New, Exon'), f = 0.9)
>
```

8 plotSamExplorer

This function generates boxplots of the metric under investigation, in our case for the number of differentially expressed genes, for different values of f .

The input argument of this function should be a `data.frame` object containing three columns with the names - `Label`, `Variable`, `Value`.

The data should look like in the following example:

Manual for the package samExploreR

```
> require(samExploreR)
> ##### Loading the example data
> data("df_sole")
> data("df_intersect")
> head(df_sole)

      Label Variable Value
1 New, Gene      0.05    3
2 New, Exon      0.05    1
3 Old, Gene      0.05    1
4 New, Gene      0.05    1
5 New, Exon      0.05    2
6 Old, Gene      0.05    0

> #head(df_intersect)
```

8.1 Plotting data using the function plotSamExplorer

Manual for the package samExploreR

```
> ### Generation of the plot:  
> require(samExploreR)  
> data("df_sole")  
> plotsamExplorer(df_sole,p.depth=.9,font.size=4, anova=FALSE,save=FALSE)
```

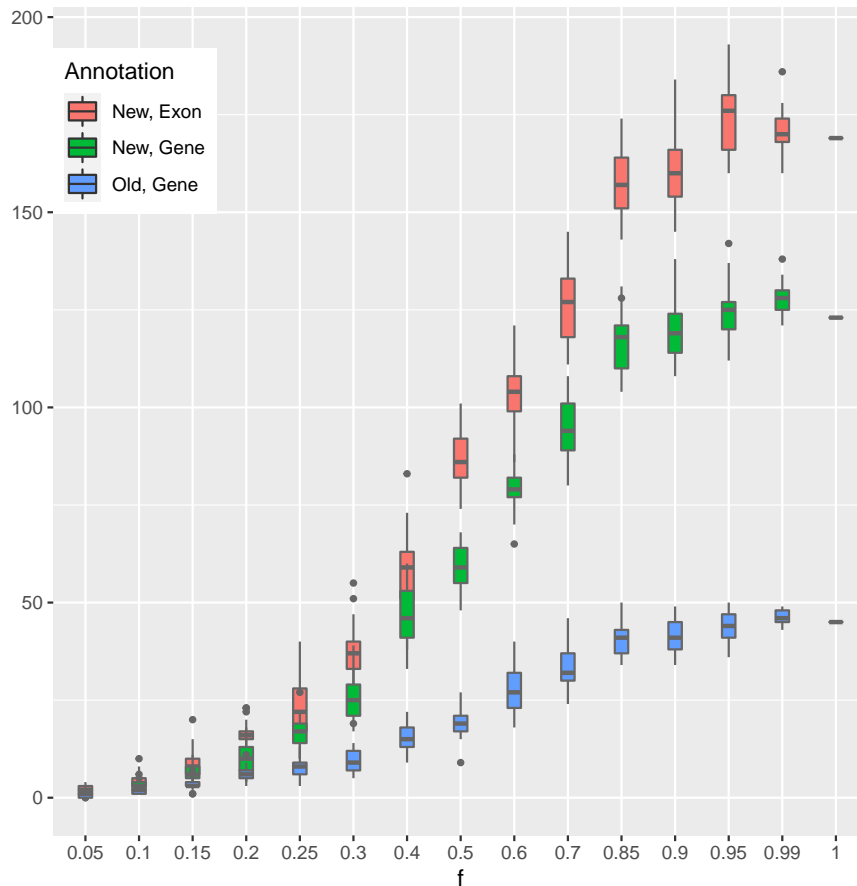


Figure 1: Boxplot of the number of differentially expressed genes for different sequence-depths f

```
> sessionInfo()
```

```
R version 4.0.0 RC (2020-04-19 r78255)  
Platform: x86_64-pc-linux-gnu (64-bit)  
Running under: Ubuntu 18.04.4 LTS
```

```
Matrix products: default
```

```
BLAS: /home/biocbuild/bbs-3.12-bioc/R/lib/libRblas.so
```

```
LAPACK: /home/biocbuild/bbs-3.12-bioc/R/lib/libRlapack.so
```

```
locale:
```

```
[1] LC_CTYPE=en_US.UTF-8
```

```
LC_NUMERIC=C
```

```
[3] LC_TIME=en_US.UTF-8
```

```
LC_COLLATE=C
```

Manual for the package samExploreR

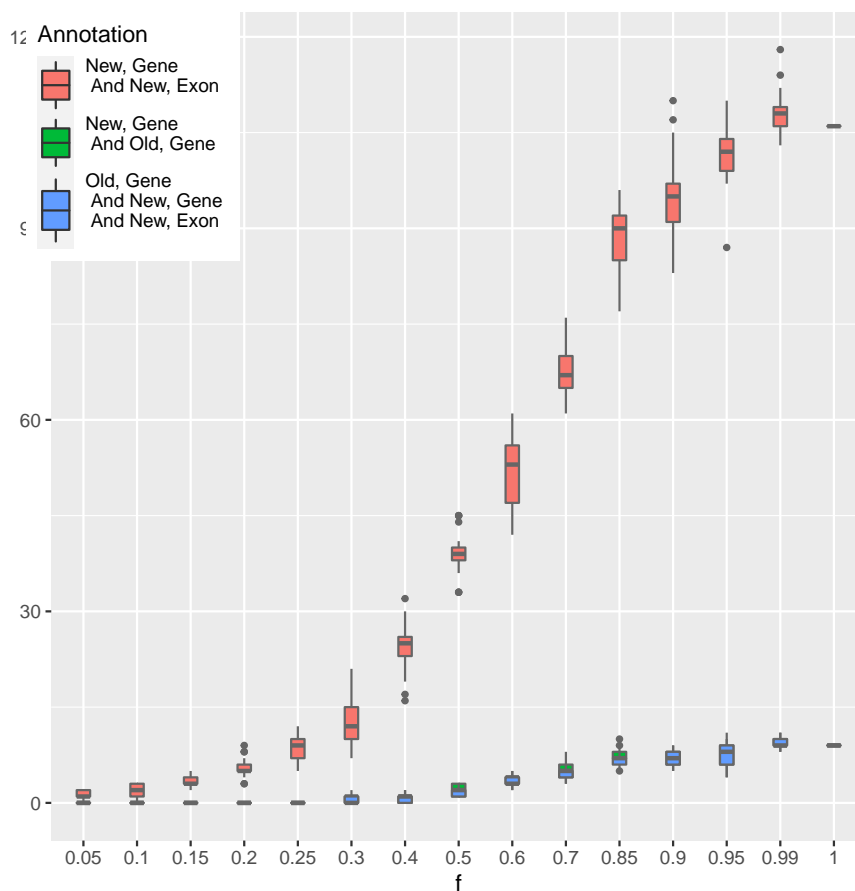


Figure 2: Boxplot of the number of differentially expressed genes for different sequence-depths f

```
[5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8 LC_NAME=C
[9] LC_ADDRESS=C LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
```

attached base packages:

```
[1] stats graphics grDevices utils datasets methods base
```

other attached packages:

```
[1] samExploreR_1.13.0 edgeR_3.31.0
[3] limma_3.45.0 RNAseqData.HNRNPC.bam.chr14_0.25.0
[5] Rsubread_2.3.0 ggplot2_3.3.0
```

loaded via a namespace (and not attached):

```
[1] Rcpp_1.0.4.6 pillar_1.4.3 compiler_4.0.0
[4] BiocManager_1.30.10 tools_4.0.0 digest_0.6.25
[7] evaluate_0.14 lifecycle_0.2.0 tibble_3.0.1
[10] gtable_0.3.0 lattice_0.20-41 pkgconfig_2.0.3
```

Manual for the package samExploreR

```
[13] rlang_0.4.5      Matrix_1.2-18    yaml_2.2.1
[16] xfun_0.13        withr_2.2.0      dplyr_0.8.5
[19] knitr_1.28       vctrs_0.2.4      locfit_1.5-9.4
[22] grid_4.0.0       tidyselect_1.0.0 glue_1.4.0
[25] R6_2.4.1         rmarkdown_2.1    farver_2.0.3
[28] purrr_0.3.4      magrittr_1.5     scales_1.1.0
[31] ellipsis_0.3.0   htmltools_0.4.0 assertthat_0.2.1
[34] BiocStyle_2.17.0 colorspace_1.4-1 labeling_0.3
[37] munsell_0.5.0    crayon_1.3.4
```

References

- [1] Alexey Stupnikov, Shailesh Tripathi, Ricardo de Matos Simoes, Darragh McArt, Manuel Salto-Tellez, Galina Glazko, Matthias Dehmer, and Frank Emmert-Streib. samExploreR: exploring reproducibility and robustness of RNA-seq results based on SAM files. *Bioinformatics*, 32(21):3345–3347, November 2016. URL: <http://dx.doi.org/10.1093/bioinformatics/btw475>, doi:10.1093/bioinformatics/btw475.
- [2] Y Liao, GK Smyth, and W Shi. featurecounts: an efficient general-purpose read summarization program. *arXiv*, 1305:16, 2013.