## Riborex Manual

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#### Introduction

Riborex is a R package for identifying differentially translated genes from Ribo-seq data. Riborex integrates both RNA- and Ribo-seq read count data into a single generalized linear model (GLM) and generates a modified design matrix reflecting the integration. At its core, Riborex applies existing RNA-seq analysis tools such as edgeR, DESeq2 and Voom to this modified design matrix and identifies differential translation across conditions.

#### Detailed example

First, we need to load Riborex library.

#### library(riborex)

```
## Warning: replacing previous import 'stats::sd' by 'BiocGenerics::sd' when
## loading 'S4Vectors'
## Warning: replacing previous import 'stats::var' by 'BiocGenerics::var' when
## loading 'S4Vectors'
## Warning: multiple methods tables found for 'var'
## Warning: multiple methods tables found for 'sd'
## Warning: multiple methods tables found for 'rowSums'
## Warning: multiple methods tables found for 'colSums'
## Warning: multiple methods tables found for 'rowMeans'
## Warning: multiple methods tables found for 'colMeans'
## Warning: replacing previous import 'stats::sd' by 'BiocGenerics::sd' when
## loading 'IRanges'
## Warning: replacing previous import 'stats::var' by 'BiocGenerics::var' when
## loading 'IRanges'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'IRanges'
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'IRanges'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'IRanges'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'IRanges'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'IRanges'
```

```
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'IRanges'
## Warning: multiple methods tables found for 'var'
## Warning: multiple methods tables found for 'sd'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'GenomeInfoDb'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'GenomicRanges'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'XVector'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'SummarizedExperiment'
```

```
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'SummarizedExperiment'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'SummarizedExperiment'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'SummarizedExperiment'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'SummarizedExperiment'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'SummarizedExperiment'
## Warning: replacing previous import 'BiocGenerics::var' by 'IRanges::var'
## when loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::sd' by 'IRanges::sd' when
## loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'DESeq2'
## Warning: replacing previous import 'BiocGenerics::rowSums' by
## 'S4Vectors::rowSums' when loading 'AnnotationDbi'
## Warning: replacing previous import 'BiocGenerics::var' by 'S4Vectors::var'
## when loading 'AnnotationDbi'
## Warning: replacing previous import 'BiocGenerics::rowMeans' by
## 'S4Vectors::rowMeans' when loading 'AnnotationDbi'
## Warning: replacing previous import 'BiocGenerics::colSums' by
## 'S4Vectors::colSums' when loading 'AnnotationDbi'
## Warning: replacing previous import 'BiocGenerics::sd' by 'S4Vectors::sd'
## when loading 'AnnotationDbi'
## Warning: replacing previous import 'BiocGenerics::colMeans' by
## 'S4Vectors::colMeans' when loading 'AnnotationDbi'
```

The input for Riborex are two read count tables summarized from RNA-seq and Ribo-seq data respectively. The read count table should be organized as a data frame with rows correspond to genes and columns correspond to samples as shown below.

```
data(riborexdata)
RNACntTable <- riborexdata$rna
RiboCntTable <- riborexdata$ribo</pre>
```

We can check the first five lines of the table:

```
head(RNACntTable,5)
```

```
##
                       BN_336 BN_337 BN_338 BN_339
                                            4
## ENSRNOGOOOOOOO017
                             7
                                   11
                                                   4
## ENSRNOG00000000024
                          2467
                                 2478
                                         3258
                                                2316
## ENSRNOG0000000033
                           206
                                  282
                                          330
                                                 244
## ENSRNOG00000000034
                           758
                                  672
                                         1335
                                                 767
## ENSRNOG0000000036
                           237
                                  163
                                          211
                                                 189
head(RiboCntTable,5)
                       BN_341 BN_342 BN_343 BN_344
##
## ENSRNOGOOOOOOO017
                            15
                                    5
                                           10
                                         2864
## ENSRNOG00000000024
                          5206
                                 5921
                                                1985
## ENSRNOG0000000033
                            30
                                   30
                                           23
                                                  13
## ENSRNOG0000000034
                           943
                                  775
                                          842
                                                 311
```

Then we need to prepare two vectors to indicate the treatments of samples in RNA- and Ribo-seq data. Both RNA-seq and Ribo-seq can have different number of samples in control and treated conditions, and RNA-seq and Ribo-seq data can have different number of samples.

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```
rnaCond <- c("control", "control", "treated", "treated")
riboCond <- c("control", "control", "treated")</pre>
```

After the two read count table and two condition vectors are ready, we can use riborex (), and we can choose which engine to use. By default, DESeq2 is used as the engine if you don't specify the engine option. Use help(riborex) in R to see more details about this function.

```
res.deseq2 <- riborex(RNACntTable, RiboCntTable, rnaCond, riboCond)</pre>
```

The format of the result is the same when DESeq2 is used in RNA-seq analysis.

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```
res.deseq2
```

## ENSRNOG0000000036

```
## log2 fold change (MLE): EXTRA1 treated vs control
## Wald test p-value: EXTRA1 treated vs control
## DataFrame with 13916 rows and 6 columns
##
                         baseMean log2FoldChange
                                                      lfcSE
                                                                   stat
##
                         <numeric>
                                        <numeric> <numeric>
                                                              <numeric>
## ENSRNOG0000000017
                         7.694934
                                       1.53323724 1.5662846
                                                              0.9789008
## ENSRNOG0000000024 3544.238071
                                      -0.10215095 0.2020511 -0.5055699
## ENSRNOG0000000033
                       111.439451
                                       0.26525296 0.5033680
                                                              0.5269564
  ENSRN0G00000000034
                       783.974916
                                       0.06513995 0.3506098
                                                              0.1857904
## ENSRNOG0000000036
                        99.151141
                                      -0.63425877 0.5910827 -1.0730457
##
                               . . .
                                              . . .
                                                         . . .
## ENSRNOG00000061895
                        105.24110
                                       0.52804379 0.4846633
                                                              1.0895065
## ENSRNOG00000061899
                         40.04057
                                      -0.49721003 0.7802896 -0.6372122
## ENSRNOG00000061910
                       4237.53481
                                       0.30904740 0.2460783
                                                              1.2558905
## ENSRNOG00000061928
                       1651.61680
                                      -0.08099135 0.2465675 -0.3284753
## ENSRNOG00000061989
                                      -0.10482910 0.4554566 -0.2301627
                        107.36281
##
                         pvalue
                                      padj
##
                       <numeric> <numeric>
## ENSRNOG0000000017 0.3276290 0.7366558
## ENSRNOG00000000024 0.6131586 0.9019205
## ENSRNOG00000000033 0.5982239 0.8963865
## ENSRNOG00000000034 0.8526091 0.9748558
  ENSRNOG00000000036 0.2832506 0.6901202
## ENSRNOG00000061895 0.2759306 0.6824115
```

```
## ENSRNOG00000061899 0.5239866 0.8635147

## ENSRNOG00000061910 0.2091557 0.5984207

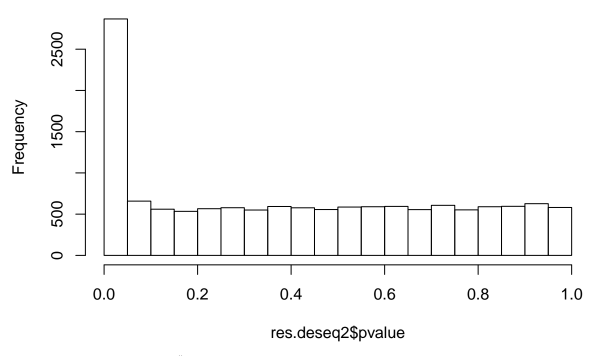
## ENSRNOG00000061928 0.7425523 0.9428604

## ENSRNOG000000061989 0.8179654 0.9676247
```

You can check the p-value distribution by

```
hist(res.deseq2$pvalue)
```

## Histogram of res.deseq2\$pvalue



Also, you can use summary () for your results.

```
summary(res.deseq2)
##
## out of 13916 with nonzero total read count
## adjusted p-value < 0.1
## LFC > 0 (up)
                    : 1107, 8%
## LFC < 0 (down)
                    : 1217, 8.7%
## outliers [1]
                    : 0, 0%
## low counts [2]
                    : 540, 3.9%
## (mean count < 6)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
And results can be saved by:
write.table(res.deseq2, "riborex_res_deseq2.txt", quote=FALSE)
```

The format of the result is the same when edgeR is used in RNA-seq analysis.

res.edgeR <- riborex(RNACntTable, RiboCntTable, rnaCond, riboCond, "edgeR")

If you want to use edgeR as your engine, you can use riborex () as:

# head(res.edgeR\$table)

```
##
                            logFC
                                     logCPM
                                                    LR
                                                          PValue
                                                                       FDR
                       1.36290149 -2.456968 1.80649827 0.1789289 0.4627345
## ENSRNOG0000000017
## ENSRNOG0000000024 -0.30127172
                                   6.212404 2.13670654 0.1438103 0.4037250
## ENSRNOG0000000033
                       0.07178854
                                   1.313235 0.02631769 0.8711269 0.9636097
## ENSRNOG00000000034 -0.13430329
                                   4.029136 0.12541035 0.7232390 0.9111612
## ENSRNOG0000000036 -0.82540899
                                   1.132478 2.15554356 0.1420562 0.4008219
## ENSRNDG0000000040 -0.19057283 -1.555003 0.07537378 0.7836675 0.9338658
```

For edgeR engine, you can also choose to estimate dispersion of RNA-seq and Ribo-seq data separately by specifying engine as "edgeRD".

```
res.edgeRD <- riborex(RNACntTable, RiboCntTable, rnaCond, riboCond, "edgeRD")
```

If you want to use Voom as the engine, you can run riborex () as:

```
res.voom <- riborex(RNACntTable, RiboCntTable, rnaCond, riboCond, "Voom")
```

The format of the result is the same when Voom is used in RNA-seq analysis.

```
head(res.voom)
```

```
##
                            logFC
                                     AveExpr
                                                          P. Value adj. P. Val
                                                      t.
                      1.39674189 -2.8437969
## ENSRNOG0000000017
                                              1.2847048 0.2268559 0.5243467
## ENSRNOG0000000024 -0.30047073
                                   6.0075571 -1.8737990 0.0893910 0.2991861
## ENSRNOG00000000033 0.07661457
                                   0.7070877
                                              0.1687028 0.8692764 0.9540355
## ENSRNOG0000000034 -0.13777833
                                   3.9701893 -0.3933633 0.7020190 0.8871606
## ENSRNOG00000000036 -0.89165405
                                   0.7106061 -1.3890222 0.1939374 0.4826701
## ENSRNOG00000000040 -0.20967655 -1.7042630 -0.2991046 0.7707701 0.9157225
##
## ENSRNOG0000000017 -4.928058
## ENSRNOG0000000024 -5.682138
## ENSRNOG00000000033 -6.320171
## ENSRNOG0000000034 -7.130133
## ENSRNOG0000000036 -5.425369
## ENSRNOG00000000040 -5.854482
```

# Multi-factor experiment

Since we don't find any available ribosome profiling data generated in a multi-factor experiement, here we generate a pseudo dataset to demonstrate the usage of riborex in a multi-factor experiment. The pseudo dataset have 8 samples in RNA-seq and Ribo-seq, and two factors are included.

```
rna <- RNACntTable[,c(1,2,3,4,1,2,3,4)]
ribo <- RiboCntTable[,c(1,2,3,4,1,2,3,4)]</pre>
```

For multi-factor experiment, we prepare two data frames to indicate the treatment under each factor. Here for the 8 samples in both RNA- and Ribo-seq experiment, the 3rd and 4th samples are treated with drug1 and the 7th and 8th samples are treated with drug2.

```
riboCond <- data.frame(factor1=(c("control1", "control1", "treated1", "treated1",
                                  "control1", "control1", "control1", "control1")),
                       factor2=(c("control2", "control2", "control2", "control2",
                                  "control2", "control2", "treated2", "treated2")))
```

Also we need to prepare a contrast to specify the comparison we want to perform, for example, if we want to

```
compare the influence of the usage of drug2. The contrast can be constructed as:
contrast = c("factor2", "control2", "treated2")
Then riborex () is used with contrast specified.
res.deseq2 <- riborex(rna, ribo, rnaCond, riboCond, "DESeq2", contrast = contrast)
We can see the summary of the result:
summary(res.deseq2)
## out of 13916 with nonzero total read count
## adjusted p-value < 0.1
## LFC > 0 (up)
                     : 1887, 14%
## LFC < 0 (down)
                     : 1987, 14%
## outliers [1]
                     : 0, 0%
## low counts [2]
                     : 270, 1.9%
## (mean count < 3)
## [1] see 'cooksCutoff' argument of ?results
## [2] see 'independentFiltering' argument of ?results
edgeR and edgeRD can be used in a similar way.
res.edgeR <- riborex(rna, ribo, rnaCond, riboCond, "edgeR", contrast = contrast)
res.edgeRD <- riborex(rna, ribo, rnaCond, riboCond, "edgeRD", contrast = contrast)</pre>
```

Currently, you can't choose Voom as the engine in a multi-factor experiment yet.

### Setup

This analysis was conducted on

```
sessionInfo()
```

```
## R version 3.3.2 (2016-10-31)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 17.04
##
## locale:
## [1] LC_CTYPE=en_US.UTF-8
                                  LC_NUMERIC=C
## [3] LC_TIME=en_US.UTF-8
                                  LC_COLLATE=en_US.UTF-8
## [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] parallel stats4
                          stats
                                    graphics grDevices utils
                                                                  datasets
```

```
## [8] methods
                 base
##
## other attached packages:
   [1] riborex_1.2.3
                                    edgeR_3.16.5
##
    [3] limma_3.30.13
                                    DESeq2_1.16.1
   [5] SummarizedExperiment_1.4.0 Biobase_2.34.0
##
   [7] GenomicRanges 1.26.4
                                    GenomeInfoDb 1.10.3
   [9] IRanges_2.8.2
                                    S4Vectors_0.12.2
##
## [11] BiocGenerics_0.22.0
##
## loaded via a namespace (and not attached):
   [1] genefilter_1.58.1
                              locfit_1.5-9.1
                                                   splines_3.3.2
##
   [4] lattice_0.20-35
                              colorspace_1.3-2
##
                                                   htmltools_0.3.6
  [7] yaml_2.1.14
                              base64enc_0.1-3
                                                   blob_1.1.0
## [10] survival_2.41-3
                             XML_3.98-1.6
                                                   rlang_0.1.2
## [13] foreign_0.8-69
                             DBI_0.7
                                                   BiocParallel_1.8.2
## [16] bit64_0.9-7
                             RColorBrewer_1.1-2
                                                   plyr_1.8.4
## [19] stringr 1.2.0
                              zlibbioc_1.20.0
                                                   munsell 0.4.3
## [22] gtable_0.2.0
                             htmlwidgets_0.9
                                                   memoise_1.1.0
## [25] evaluate_0.10.1
                              latticeExtra_0.6-28
                                                   knitr 1.17
## [28]
       geneplotter_1.52.0
                             AnnotationDbi_1.38.0
                                                   htmlTable_1.9
## [31] Rcpp_0.12.12
                              acepack_1.4.1
                                                   xtable_1.8-2
## [34] scales_0.5.0
                                                   checkmate_1.8.3
                              backports_1.1.0
## [37] Hmisc 4.0-3
                              annotate_1.52.1
                                                   XVector_0.14.1
## [40] bit_1.1-12
                              gridExtra_2.3
                                                   ggplot2_2.2.1
## [43] digest_0.6.12
                              stringi_1.1.5
                                                   grid_3.3.2
## [46] rprojroot_1.2
                              tools_3.3.2
                                                   bitops_1.0-6
                              lazyeval_0.2.0
                                                   RCurl_1.95-4.8
## [49]
       magrittr_1.5
## [52] tibble_1.3.4
                             RSQLite_2.0
                                                   Formula_1.2-2
## [55] cluster_2.0.6
                             Matrix_1.2-11
                                                   data.table_1.10.4
## [58] rmarkdown_1.6
                             rpart_4.1-11
                                                   nnet_7.3-12
```