

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)
```

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
```

We can check the first five lines of the table:

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

##	BN_336	BN_337	BN_338	BN_339
## ENSRNOG000000000017	7	11	4	4
## ENSRNOG000000000024	2467	2478	3258	2316
## ENSRNOG000000000033	206	282	330	244
## ENSRNOG000000000034	758	672	1335	767
## ENSRNOG000000000036	237	163	211	189

```
head(RiboCntTable,5)
```

##	BN_341	BN_342	BN_343	BN_344
## ENSRNOG000000000017	15	5	10	2
## ENSRNOG000000000024	5206	5921	2864	1985
## ENSRNOG000000000033	30	30	23	13
## ENSRNOG000000000034	943	775	842	311
## ENSRNOG000000000036	80	49	30	7

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.

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

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)
```

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

```
res.deseq2
```

```
## log2 fold change (MAP): 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>
## ENSRNOG000000000017	7.694934	0.60242172	0.6633528	0.9081467
## ENSRNOG000000000024	3544.238071	-0.09262781	0.1967739	-0.4707321
## ENSRNOG000000000033	111.439451	0.12090513	0.4390872	0.2753556
## ENSRNOG000000000034	783.974916	0.07400898	0.3243979	0.2281426
## ENSRNOG000000000036	99.151141	-0.65373460	0.4894467	-1.3356604
##
## ENSRNOG000000061895	105.24110	0.42802240	0.4224309	1.0132366
## ENSRNOG000000061899	40.04057	-0.55910038	0.5828614	-0.9592339
## ENSRNOG000000061910	4237.53481	0.29957699	0.2366771	1.2657623
## ENSRNOG000000061928	1651.61680	-0.07583546	0.2370871	-0.3198633
## ENSRNOG000000061989	107.36281	-0.06620802	0.4009571	-0.1651250
##	pvalue	padj		
##	<numeric>	<numeric>		
## ENSRNOG000000000017	0.3638007	0.7494602		
## ENSRNOG000000000024	0.6378320	0.9022741		
## ENSRNOG000000000033	0.7830430	0.9509349		
## ENSRNOG000000000034	0.8195354	0.9607316		
## ENSRNOG000000000036	0.1816603	0.5432690		
##		
## ENSRNOG000000061895	0.3109472	0.7034611		
## ENSRNOG000000061899	0.3374409	0.7261818		
## ENSRNOG000000061910	0.2055982	0.5775201		
## ENSRNOG000000061928	0.7490719	0.9411896		
## ENSRNOG000000061989	0.8688456	0.9744379		

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)      : 1139, 8.2%
## LFC < 0 (down)    : 1274, 9.2%
## 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
```

And results can be saved by:

```
write.table(res.deseq2, "riborex_res_deseq2.txt", quote=FALSE)
```

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

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

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

```
head(res.edgeR$table)
```

```
##              logFC    logCPM      LR    PValue      FDR
## ENSRNOG000000000017  1.3629124 -2.456968 1.80670421 0.1789041 0.4626704
## ENSRNOG000000000024 -0.3012715  6.212404 2.13615823 0.1438618 0.4037879
## ENSRNOG000000000033  0.0717875  1.313235 0.02631400 0.8711358 0.9636269
## ENSRNOG000000000034 -0.1343023  4.029136 0.12544284 0.7232047 0.9111520
## ENSRNOG000000000036 -0.8253990  1.132478 2.15594580 0.1420190 0.4007169
## ENSRNOG000000000040 -0.1905669 -1.555003 0.07536038 0.7836862 0.9338705
```

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      t    P.Value adj.P.Val
## ENSRNOG000000000017  1.39674189 -2.8437969  1.2847048 0.2268559 0.5243467
## ENSRNOG000000000024 -0.30047073  6.0075571 -1.8737990 0.0893910 0.2991861
## ENSRNOG000000000033  0.07661457  0.7070877  0.1687028 0.8692764 0.9540355
## ENSRNOG000000000034 -0.13777833  3.9701893 -0.3933633 0.7020190 0.8871606
## ENSRNOG000000000036 -0.89165405  0.7106061 -1.3890222 0.1939374 0.4826701
## ENSRNOG000000000040 -0.20967655 -1.7042630 -0.2991046 0.7707701 0.9157225
```

```
##                                     B
## ENSRNOG00000000017 -4.928058
## ENSRNOG00000000024 -5.682138
## ENSRNOG00000000033 -6.320171
## ENSRNOG00000000034 -7.130133
## ENSRNOG00000000036 -5.425369
## ENSRNOG00000000040 -5.854482
```

Multi-factor experiment

Since we don't find any available ribosome profiling data generated in a multi-factor experiment, 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)]
```

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.

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

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)      : 1889, 14%
## LFC < 0 (down)    : 1972, 14%
## outliers [1]      : 0, 0%
```

```
## low counts [2] : 810, 5.8%
## (mean count < 8)
## [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)
```

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

Setup

This analysis was conducted on

```
sessionInfo()
```

```
## R version 3.3.1 (2016-06-21)
## Platform: x86_64-pc-linux-gnu (64-bit)
## Running under: Ubuntu 14.04.4 LTS
##
## 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_1.0.0             edgeR_3.14.0
##  [3] limma_3.28.11            DESeq2_1.12.3
##  [5] SummarizedExperiment_1.2.3 Biobase_2.32.0
##  [7] GenomicRanges_1.24.2     GenomeInfoDb_1.8.1
##  [9] IRanges_2.6.1            S4Vectors_0.10.1
## [11] BiocGenerics_0.18.0
##
## loaded via a namespace (and not attached):
##  [1] Rcpp_0.12.5              formatR_1.4              RColorBrewer_1.1-2
##  [4] plyr_1.8.4               XVector_0.12.0           tools_3.3.1
##  [7] zlibbioc_1.18.0          rpart_4.1-10            digest_0.6.9
## [10] RSQLite_1.0.0            annotate_1.50.0          evaluate_0.9
## [13] gtable_0.2.0             lattice_0.20-33         Matrix_1.2-6
## [16] DBI_0.4-1                yaml_2.1.13             gridExtra_2.2.1
## [19] genefilter_1.54.2        stringr_1.0.0           knitr_1.13
```

## [22] cluster_2.0.4	locfit_1.5-9.1	grid_3.3.1
## [25] nnet_7.3-12	data.table_1.9.6	AnnotationDbi_1.34.3
## [28] XML_3.98-1.4	survival_2.39-4	BiocParallel_1.6.2
## [31] foreign_0.8-66	rmarkdown_1.0	latticeExtra_0.6-28
## [34] Formula_1.2-1	geneplotter_1.50.0	ggplot2_2.1.0
## [37] magrittr_1.5	Hmisc_3.17-4	scales_0.4.0
## [40] htmltools_0.3.5	splines_3.3.1	xtable_1.8-2
## [43] colorspace_1.2-6	stringi_1.1.1	acepack_1.3-3.3
## [46] munsell_0.4.3	chron_2.3-47	