Pre-Processing for the Zebrafish RNA-Seq Gene-Level Counts

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This vignette describes the pre-processing steps that were followed for the generation of the gene-level read counts contained in the *Bioconductor* package *zebrafishRNASeq*.

Contents

1	Sample preparation and sequencing.	1
2	Read alignment and expression quantitation	1
3	Loading the zebrafish data into R	2
4	Session info	2

1 Sample preparation and sequencing

Olfactory sensory neurons were isolated from three pairs of gallein-treated and control embryonic zebrafish pools and purified by fluorescence activated cell sorting (FACS) [1]. Each RNA sample was enriched in poly(A)+ RNA from 10–30 ng total RNA and 1 μ L (1:1000 dilution) of Ambion ERCC ExFold RNA Spike-in Control Mix 1 was added to 30 ng of total RNA before mRNA isolation. cDNA libraries were prepared according to manufacturer's protocol. The six libraries were sequenced in two multiplex runs on an Illumina HiSeq2000 sequencer, yielding approximately 50 million 100bp paired-end reads per library.

2 Read alignment and expression quantitation

We made use of a custom reference sequence, defined as the union of the zebrafish reference genome (Zv9, downloaded from Ensembl [2], v. 67) and the ERCC spike-in sequences (http://tools.invitrogen.com/downloads/ERCC92.fa). Reads were mapped with TopHat [3] (v. 2.0.4), with the following parameters,

--library-type=fr-unstranded -G ensembl.gtf --transcriptome-index=transcript --no-novel-juncs where ensembl.gtf is a GTF file containing Ensembl gene annotation.

Gene-level read counts were obtained using the htseq-count python script [4] in the "union" mode and Ensembl (v. 67) gene annotation.

Pre-Processing for the Zebrafish RNA-Seq Gene-Level Counts

After verifying that there were no run-specific biases, we used the sums of the counts of the two runs as the expression measures for each library.

3 Loading the zebrafish data into *R*

To load the gene-level read counts into R, simply type

```
library(zebrafishRNASeq)
data(zfGenes)
head(zfGenes)
                      Ctl1 Ctl3 Ctl5 Trt9 Trt11 Trt13
## ENSDARG0000000001
                      304 129
                                339
                                      102
                                             16
                                                  617
## ENSDARG00000000002
                      605
                            637
                                 406
                                       82
                                            230
                                                 1245
## ENSDARG0000000018 391
                            235
                                                  565
                                217
                                      554
                                            451
## ENSDARG0000000019 2979 4729 7002 7309
                                           9395
                                                 3349
## ENSDARG00000000068
                       89
                            356
                                  41
                                      149
                                             45
                                                   44
## ENSDARG00000000009 312 184
                                 844
                                      269
                                            513
                                                  243
```

The ERCC spike-in read counts are in the last rows of the same matrix and can be retrieved in the following way.

```
spikes <- zfGenes[grep("^ERCC", rownames(zfGenes)),]</pre>
head(spikes)
##
                Ctl1
                       Ctl3
                             Ctl5
                                     Trt9 Trt11 Trt13
## ERCC-00002 97227 38556
                             68367 148331 169360 100974
## ERCC-00003 10925
                       6240 11156 36652 21184 21841
## ERCC-00004 379182 179870 256130 679783 529085 311169
## ERCC-00009
                2452
                       1183
                              1042
                                      1895
                                             3520
## ERCC-00012
                   0
                          0
                                 0
                                        0
                                                0
                                                       0
## ERCC-00013
                                       205
                                               21
                                                       3
                  89
                          8
                                 0
```

The typical use of this dataset is the indentification of differentially expressed genes between control (Ctl) and treated (Trt) samples. For additional details, exploratory analysis, and normalization of the zebrafish data see [5, 6]. The data are used as a case study for the *Bioconductor* package *RUVSeq*.

4 Session info

```
toLatex(sessionInfo())
```

- R version 3.5.1 Patched (2018-07-12 r74967), 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 16.04.5 LTS

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- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.8-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.8-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: zebrafishRNASeq 1.2.0
- Loaded via a namespace (and not attached): BiocManager 1.30.3, BiocStyle 2.10.0, Rcpp 0.12.19, backports 1.1.2, compiler 3.5.1, digest 0.6.18, evaluate 0.12, highr 0.7, htmltools 0.3.6, knitr 1.20, magrittr 1.5, rmarkdown 1.10, rprojroot 1.3-2, stringi 1.2.4, stringr 1.3.1, tools 3.5.1, yaml 2.2.0

References

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- [2] P. Flicek, M. R. Amode, D. Barrell, K. Beal, S. Brent, D. Carvalho-Silva, P. Clapham, G. Coates, S. Fairley, S. Fitzgerald, et al. Ensembl 2012. *Nucleic Acids Research*, 40(D1):D84–D90, 2012.
- [3] C. Trapnell, L. Pachter, and S. L. Salzberg. TopHat: discovering splice junctions with RNA-Seq. *Bioinformatics*, 25(9):1105–1111, 2009.
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- [5] D. Risso, J. Ngai, T.P. Speed, and S. Dudoit. Using controls for the normalization of RNA-Seq data. *Nature Biotechnology*, 2014. Accepted.
- [6] D. Risso, J. Ngai, T.P. Speed, and S. Dudoit. The role of spike-in standards in the normalization of RNA-seq. In D. Nettleton and S. Datta, editors, *Statistical Analysis of Next Generation Sequence Data*. Springer, 2014.