# Gene set enrichment analysis of RNA-Seq data with the SeqGSEA package

Xi Wang $^{1,2}$  and Murray Cairns $^{1,2,3}$ 

October 30, 2017

<sup>1</sup>School of Biomedical Sciences and Pharmacy, The University of Newcastle, Callaghan, New South Wales, Australia

#### xi.wang@mdc-berlin.de

## Contents

| 1        | Introduction                                 |   |    |  |  |  |  |  |  |
|----------|--|---|----|--|--|--|--|--|--|
|          | 1.1  | Background                                      | 2  |  |  |  |  |  |  |
|          | 1.2  | Getting started                                 | 2  |  |  |  |  |  |  |
|          | 1.3  | Package citation                                |    |  |  |  |  |  |  |
| <b>2</b> | Differential splicing analysis and DS scores |   |    |  |  |  |  |  |  |
| _        | 2.1  | The ReadCountSet class                          | :  |  |  |  |  |  |  |
|          | 2.2  | DS analysis and DS scores                       |    |  |  |  |  |  |  |
|          | 2.3  | DS permutation p-values                         |    |  |  |  |  |  |  |
|          | ۷.5  | D5 permutation p-values                         | و  |  |  |  |  |  |  |
| 3        | Diff   | ferential expression analysis and DE scores     | 6  |  |  |  |  |  |  |
|          | 3.1  | Gene read count data from ReadCountSet class    | (  |  |  |  |  |  |  |
|          | 3.2  | DE analysis and DE scores                       | 6  |  |  |  |  |  |  |
|          | 3.3  | DE permutation p-values                         | 8  |  |  |  |  |  |  |
| 4        | Integrative GSEA runs                        |   |    |  |  |  |  |  |  |
|          | 4.1  | DE/DS score integration                         | 8  |  |  |  |  |  |  |
|          | 4.2  | Initialization of SegGeneSet objects            | (  |  |  |  |  |  |  |
|          | 4.3  | running GSEA with integrated gene scores        |    |  |  |  |  |  |  |
|          | 4.4  | SeqGSEA result displays                         | 12 |  |  |  |  |  |  |
|          | 1.1  | Soq OS 211 Too all all plays                    |    |  |  |  |  |  |  |
| 5        | Rui  | nning SeqGSEA with multiple cores               | 13 |  |  |  |  |  |  |
|          | 5.1  | R-parallel packages                             | 13 |  |  |  |  |  |  |
|          | 5.2  | Parallelizing analysis on permutation data sets | 15 |  |  |  |  |  |  |
| 6        | Analysis examples 1                          |   |    |  |  |  |  |  |  |
|          | 6.1  | Starting from your own RNA-Seq data             | 15 |  |  |  |  |  |  |
|          | 6.2  | Exemplified pipeline for integrating DE and DS  |    |  |  |  |  |  |  |
|          | 6.3  | Exemplified pipeline for DE-only analysis       | 17 |  |  |  |  |  |  |
|          | 6.4  | One-step SeqGSEA analysis                       | 18 |  |  |  |  |  |  |

 $<sup>^2\</sup>mathrm{Hunter}$  Medical Research Institute, New Lambton, New South Wales, Australia

 $<sup>^3{\</sup>rm Schizophrenia}$ Research Institute, Sydney, New South Wales, Australia

7 Session information 19

#### 1 Introduction

#### 1.1 Background

Transcriptome sequencing (RNA-Seq) has become a key technology in transcriptome studies because it can quantify overall expression levels and the degree of alternative splicing for each gene simultaneously. Many methods and tools, including quite a few R/Bioconductor packages, have been developed to deal with RNA-Seq data for differential expression analysis and thereafter functional analysis aiming at novel biological and biomedical discoveries. However, those tools mainly focus on each gene's overall expression and may miss the opportunities for discoveries regarding alternative splicing or the combination of the two.

SeqGSEA is novel R/Bioconductor package to derive biological insight by integrating differential expression (DE) and differential splicing (DS) from RNA-Seq data with functional gene set analysis. Due to the digital feature of RNA-Seq count data, the package utilizes negative binomial distributions for statistical modeling to first score differential expression and splicing in each gene, respectively. Then, integration strategies are applied to combine the two scores for integrated gene set enrichment analysis. See the publications Wang and Cairns (2013) and Wang and Cairns (2014) for more details. The SeqGSEA package can also give detection results of differentially expressed genes and differentially spliced genes based on sample label permutation.

#### 1.2 Getting started

The SeqGSEA depends on Biobase for definitions of class ReadCountSet and class SeqGeneSet, DESeq for differential expression analysis, biomaRt for gene IDs/names conversion, and doParallel for parallelizing jobs to reduce running time. Make sure you have these dependent packages installed before you install SeqGSEA.

To load the SeqGSEA package, type library(SeqGSEA). To get an overview of this package, type ?SeqGSEA.

> library(SeqGSEA)

#### > ? SeqGSEA

In this Users' Guide of the SeqGSEA package, an analysis example is given in Section 6, and detailed guides for DE, DS, and integrative GSEA analysis are given in Sections 3, 2, and 4, respectively. A guide to parallelize those analyses is given in Section 5.

#### 1.3 Package citation

To cite this package, please cite the article below:

Wang X and Cairns MJ (2014). SeqGSEA: a Bioconductor package for gene set enrichment analysis of RNA-Seq data integrating differential expression and splicing. *Bioinformatics*, 30(12):1777-9.

To cite/discuss the method used in this package, please cite the article below:

Wang X and Cairns MJ (2013). Gene set enrichment analysis of RNA-Seq data: integrating differential expression and splicing. *BMC Bioinformatics*, 14(Suppl 5):S16.

# 2 Differential splicing analysis and DS scores

#### 2.1 The ReadCountSet class

To facilitate differential splicing (DS) analysis, SeqGSEA saves exon read count data using Read-CountSet class, which is derived from eSet. While below is an example showing the steps to create a new ReadCountSet object, creating a ReadCountSet object from your own data should refer to Section 6.

```
> rcounts <- cbind(t(sapply(1:10, function(x) {rnbinom(5, size=10, prob=runif(1))} )),</pre>
                   t(sapply(1:10, function(x) {rnbinom(5, size=10, prob=runif(1))} )))
> colnames(rcounts) <- c(paste("S", 1:5, sep=""), paste("C", 1:5, sep=""))</pre>
> geneIDs <- c(rep("G1", 4), rep("G2", 6))
> exonIDs <- c(paste("E", 1:4, sep=""), paste("E", 1:6, sep=""))
> RCS <- newReadCountSet(rcounts, exonIDs, geneIDs)
> RCS
ReadCountSet (storageMode: environment)
assayData: 10 features, 10 samples
  element names: counts
protocolData: none
phenoData
  sampleNames: S1 S2 ... C5 (10 total)
 varLabels: label
 varMetadata: labelDescription
featureData
 featureNames: 1 2 ... 10 (10 total)
 fvarLabels: exonIDs geneIDs ... padjust (10 total)
  fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

## 2.2 DS analysis and DS scores

To better illustrate DS analysis functions, we load an example ReadCountSet object from a real RNA-Seq data set as follows.

```
> data(RCS_example, package="SeqGSEA")
> RCS_example
ReadCountSet (storageMode: environment)
assayData: 5000 features, 20 samples
  element names: counts
protocolData: none
phenoData
 sampleNames: S1 S2 ... C10 (20 total)
 varLabels: label
 varMetadata: labelDescription
featureData
  featureNames: ENSG00000000003:001 ENSG0000000003:002 ...
    ENSG0000007402:038 (5000 total)
 fvarLabels: exonIDs geneIDs ... padjust (10 total)
 fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

This example ReadCountSet object is comprised of 20 samples and 5,000 exons, part of the prostate cancer RNA-Seq data set (Kannan et al., 2011). With the function geneID and the script below, we can easily check the number of genes involved in this data set.

#### > length(unique(geneID(RCS\_example)))

#### [1] 182

Noticed that some exons are too short or not expressed, we should first filter out these exons from following analysis to secure the robustness of our analysis. By default, function exonTestablity marks exons with the sum of read counts across all samples less than cutoff (default: 5) to be excluded in downstream analysis. Users can also exclude genes with no or low expression from downstream analysis by checking geneTestability.

```
> RCS_example <- exonTestability(RCS_example, cutoff = 5)
```

Then, the main DS analysis is executed using function estiExonNBstat for exon DS NB-statistics and function estiGeneNBstat for gene DS NB-statistics by averaging exon NB-statistics. Please refer to Wang et al. (2013) for detailed statistic analysis regarding differential splicing from exon count data.

```
> RCS_example <- estiExonNBstat(RCS_example)
> RCS_example <- estiGeneNBstat(RCS_example)
> head(fData(RCS_example)[, c("exonIDs", "geneIDs", "testable", "NBstat")])
```

|                     | exonIDs | geneIDs        | testable | NBstat    |
|---------------------|---------|----------------|----------|-----------|
| ENSG0000000003:001  | E001    | ENSG0000000003 | TRUE     | 2.0219857 |
| ENSG0000000003:002  | E002    | ENSG0000000003 | TRUE     | 0.2486443 |
| ENSG0000000003:003  | E003    | ENSG0000000003 | TRUE     | 0.1238136 |
| ENSG00000000003:004 | E004    | ENSG0000000003 | TRUE     | 1.2058520 |
| ENSG0000000003:005  | E005    | ENSG0000000003 | TRUE     | 2.0668287 |
| ENSG0000000003:006  | E006    | ENSG0000000003 | TRUE     | 0.2678247 |

We run DS analysis on the permutation data sets as well. Here we set to run permutation 20 times for demonstration; however, in practice at least 1,000 permutations are recommended. To do so, we first generate a permutation matrix, each column corresponding to each permutation; then run DS analysis on the permutation data sets, and updated permute\_NBstat\_gene slot for results.

- > permuteMat <- genpermuteMat(RCS\_example, times=20)</pre>
- > RCS\_example <- DSpermute4GSEA(RCS\_example, permuteMat)
- > head(RCS\_example@permute\_NBstat\_gene)

```
result.1 result.2 result.3 result.4 result.5 result.6
ENSG00000000003 0.8324813 1.1065575 0.4608722 0.6762978 1.7100655 0.4265713
ENSG000000000005 0.3948804 0.8202932 1.2094351 0.5989295 0.6835109 0.8423347
ENSG000000000419 0.4866245 1.8242975 0.5213321 0.6368590 1.7821468 0.9315022
ENSG000000000457 1.1720663 0.9755041 0.8341588 0.5315989 1.6485482 0.6066783
ENSG000000000460 0.7120520 0.8511214 0.6209427 0.5616192 0.6687020 0.8453076
ENSG000000000938 0.8713850 1.2656447 1.1735832 1.4138664 1.0293263 1.3163994
result.7 result.8 result.9 result.10 result.11 result.12
ENSG00000000000 0.9517230 1.0719824 0.7216319 0.3243202 0.2433626 1.0923860
ENSG000000000457 0.6863151 1.0325120 0.5653264 1.5999645 0.8013277 0.4751253
ENSG000000000038 1.8938594 0.9043653 0.9808110 1.5481539 1.5096625 0.9453188
```

```
Pesult.13 result.14 result.15 result.16 result.17 result.18 ENSG00000000000 0.6966410 0.8493780 0.9754674 0.9036558 0.2178080 0.5015296 ENSG00000000005 0.9838755 0.5124798 1.1914483 0.4790461 0.7216319 0.5934397 ENSG00000000419 0.6738308 0.8265125 0.9297117 0.6393698 0.5103505 0.6861051 ENSG00000000457 1.1382310 0.4115860 0.6561267 0.9473009 0.9385539 0.9276930 ENSG00000000460 0.8091019 0.6828277 0.4486701 0.8305067 0.7170622 0.5667417 ENSG0000000003 0.8984333 0.9713267 1.0273000 1.2470196 1.2849111 1.3969478 result.19 result.20 ENSG00000000003 1.5316286 1.7056188 ENSG00000000001 0.6703627 0.2019705 ENSG00000000457 0.5162404 0.5703134 ENSG00000000460 0.5611028 1.1170882 ENSG00000000038 1.0583110 0.7388838
```

The DS NB-statistics from the permutation data sets offer an empirical background of NB-statistics on the real data set. By normalizing NB-statistics against this background, we get the DS scores, which will be used in integrated GSEA runs (Section 4).

```
> DSscore.normFac <- normFactor(RCS_example@permute_NBstat_gene)
> DSscore <- scoreNormalization(RCS_example@featureData_gene$NBstat,
                                DSscore.normFac)
> DSscore.perm <- scoreNormalization(RCS_example@permute_NBstat_gene,
                                     DSscore.normFac)
> DSscore[1:5]
ENSG0000000003 ENSG0000000005 ENSG00000000419 ENSG00000000457 ENSG00000000460
      1.6149004
                                     1.7961059
                     0.6379990
                                                     3.4157597
                                                                     0.9984133
> DSscore.perm[1:5,1:10]
                result.1 result.2 result.3 result.4 result.5 result.6
ENSG0000000000 0.9325544 1.239577 0.5162739 0.7575960 1.9156335 0.4778497
ENSG00000000005 0.5519312 1.146538 1.6904484 0.8371342 0.9553551 1.1773459
ENSG00000000419 0.5190385 1.945814 0.5560580 0.6792800 1.9008554 0.9935494
ENSG00000000457 1.3760546 1.145282 0.9793372 0.6241193 1.9354642 0.7122656
ENSG00000000460 0.9465507 1.131420 0.8254366 0.7465761 0.8889244 1.1236912
                result.7 result.8 result.9 result.10
ENSG00000000003 2.5088743 0.6359584 0.9132714 0.4797477
ENSG0000000005 1.3302398 1.4983284 1.0086374 0.4533080
ENSG00000000419 3.3753386 1.0393987 0.4394762 0.8933049
ENSG00000000457 0.8057625 1.2122121 0.6637167 1.8784250
ENSG00000000460 1.1799963 1.0679628 1.6157099 0.9732735
```

## 2.3 DS permutation p-values

Besides calculating DS scores, based on the NB statistics on the real data set and the permutation data sets, we can also calculate a permutation p-value for each gene's DS significance in the studied data set.

```
2 ENSG00000000005 0.4564578 0.80 0.8138728
3 ENSG00000000419 1.6839390 0.15 0.2295652
4 ENSG00000000457 2.9094026 0.00 0.0000000
5 ENSG00000000460 0.7510661 0.45 0.5044586
6 ENSG000000000938 1.7949049 0.05 0.1035294
```

The adjusted p-values accounting for multiple testings were given by the BH method (Benjamini and Hochberg, 1995). Users can also apply function topDSGenes and function topDSExons to quickly get the most significant DS genes and exons, respectively.

## 3 Differential expression analysis and DE scores

#### 3.1 Gene read count data from ReadCountSet class

For gene DE analysis, read counts on each gene should be first calculated. With SeqGSEA, users usually analyze DE and DS simultaneously, so the package includes the function getGeneCount to facilitate gene read count calculation from a ReadCountSet object.

```
> geneCounts <- getGeneCount(RCS_example)
> dim(geneCounts) # 182
[1] 182 20
> head(geneCounts)
                     S2
                         S3
                              S4
                                  S5
                                      S6
                                           S7
                                                S8
                                                    S9 S10
                                                            C1 C2
                                                                    СЗ
                                                                         C4
                 S1
ENSG00000000003 495 235 386 272 255 815 1065 803 839 885 278 270 238 175 292
ENSG0000000005
                 19
                      1
                           0
                               2
                                   2
                                     12
                                            7
                                                 3
                                                     1
                                                         4
                                                             4
                                                                 4
                                                                     2
ENSG00000000419 196 134 165 184 132 344
                                          343 307 342 280 179 156 100 120 126
                                                                             52
ENSG00000000457
                     78 141
                              72 102 219
                                          344 277 337 249
                                                            62
                                                                48
                                                                     40
                                                                         43
                 97
                              25
ENSG00000000460
                 52
                      35
                          48
                                  47 105
                                          124
                                                80 156 145
                                                            48
                                                                36
                                                                     21
                                                                         34
                                                                             19
ENSG00000000938
                 27
                      44
                          57
                              43
                                  14
                                      71
                                           74 146 148 165
                                                            48
                                                                59
                                                                     32
                                                                         79
                                                                             20
                 C6
                      C7
                           C8
                              C9 C10
ENSG0000000003 432
                     519 621 475 560
ENSG0000000005
                  9
                       3
                            1 14
ENSG0000000419 169
                     255 171 164 201
ENSG0000000457 170
                     165 131 183 185
ENSG0000000460 68
                      90 48
                              72
                                   38
ENSG00000000938 103 1285 137 156
```

This function results in a matrix of 182 rows and 20 columns, corresponding to 182 genes and 20 samples.

#### 3.2 DE analysis and DE scores

DE analysis has been implemented in several R/Bioconductor packages, of which *DESeq* (Anders and Huber, 2010) is mainly utilized in *SeqGSEA* for DE analysis. With *DESeq*, we can model count data with negative binomial distributions for accounting biological variations and various biases introduced in RNA-Seq. Given the read count data on individual genes and sample grouping information, basic DE analysis based on *DESeq* including size factor estimation and dispersion estimation, is encapsulated in the function runDESeq.

```
> label <- label(RCS_example)
> DEG <- runDESeq(geneCounts, label)</pre>
```

The function runDESeq returns a CountDataSet object, which is defined in the *DESeq* package. The DE analysis in the *DESeq* package continues with the output CountDataSet object and conducts negative-binomial-based statistical tests for DE genes (using nbinomTest or nbinomGLMTest). However, in this SeqGSEA package, we define NB statistics to quantify each gene's expression difference between sample groups.

The NB statistics for DE can be achieved by the following scripts.

```
> DEGres <- DENBStat4GSEA(DEG)
> DEGres[1:5, "NBstat"]
```

```
[1] 0.5426504 0.2503510 0.0231052 14.3384053 1.4101270
```

Similarly, we run DE analysis on the permutation data sets as well. The permuteMat should be the same as used in DS analysis on the permutation data sets.

```
> DEpermNBstat <- DENBStatPermut4GSEA(DEG, permuteMat)
```

> DEpermNBstat[1:5, 1:10]

```
result.1
                   result.2
                               result.3
                                           result.4
                                                        result.5
                                                                     result.6
[1,] 3.35421107 0.740741893 1.73002781 0.16150086 9.678992631 0.052829418
[2,] 0.04940538 0.004797275 0.26701914 2.99358336 0.003209478 0.016508353
[3,] 1.44802757 0.696290373 0.09210627 1.90602747 3.085479106 0.164276441
[4,] \quad 0.35780419 \quad 0.097334542 \quad 0.62864123 \quad 0.13245463 \quad 0.633131350 \quad 0.024174850
[5,] 0.32940359 1.539203091 2.23796218 0.07354275 0.743135344 0.001517188
       result.7
                    result.8 result.9 result.10
[1,] 5.85408842 6.871886e-01 0.2538945 0.03670422
[2,] 0.08880201 7.572803e-02 0.0701655 0.08759372
[3,] 6.56900410 4.997894e-01 0.4278152 0.70213436
[4,] 0.50816382 6.167627e-06 0.3961494 0.70186665
[5,] 2.46775645 1.019120e-01 2.3086999 1.48961747
```

Once again, the DE NB-statistics from the permutation data sets offer an empirical background, so we can normalize NB-statistics against this background. By doing so, we get the DE scores, which will also be used in integrated GSEA runs (Section 4).

```
> DEscore.normFac <- normFactor(DEpermNBstat)</pre>
> DEscore <- scoreNormalization(DEGres$NBstat, DEscore.normFac)
> DEscore.perm <- scoreNormalization(DEpermNBstat, DEscore.normFac)
> DEscore[1:5]
    0.33030546 0.32218059 0.02192531 24.56948027
> DEscore.perm[1:5, 1:10]
                   result.2
                              result.3
                                         result.4
                                                     result.5
       result.1
                                                                  result.6
[1,] 2.04167226 0.450881637 1.05304935 0.09830384 5.891498891 0.032156700
[2,] 0.06358055 0.006173688 0.34363110 3.85248912 0.004130327 0.021244857
[3,] 1.37408237 0.660733504 0.08740275 1.80869399 2.927915565 0.155887476
[4,] 0.61311302 0.166786966 1.07720405 0.22696676 1.084898067 0.041424655
[5,] 0.29954227 1.399670224 2.03508494 0.06687591 0.675768143 0.001379651
      result.7
                   result.8
                              result.9 result.10
[1,] 3.5633208 0.4182843075 0.15454287 0.02234147
[2,] 0.1142807 0.0974555815 0.09029708 0.11272572
[3,] 6.2335503 0.4742671247 0.40596830 0.66627906
[4,] 0.8707608 0.0000105685 0.67881923 1.20267899
[5,] 2.2440477 0.0926734119 2.09941014 1.35457968
```

#### 3.3 DE permutation p-values

Similar to DS analysis, comparing NB-statistics on the real data set and those on the permutation data sets, we can get permutation p-values for each gene's DE significance.

```
> DEGres <- DEpermutePval(DEGres, DEpermNBstat)
> DEGres[1:6, c("NBstat", "perm.pval", "perm.padj")]
```

```
NBstat perm.pval perm.padj
ENSG0000000003
                 0.5426504
                                 0.55
ENSG00000000005
                 0.2503510
                                 0.30
                                               1
                                 0.95
ENSG00000000419
                 0.0231052
                                               1
ENSG0000000457 14.3384053
                                 0.00
                                               0
ENSG00000000460
                 1.4101270
                                 0.40
                                               1
                 2.1976989
                                 0.00
                                               0
ENSG00000000938
```

For a comparison to the nominal p-values from exact testing and forming comprehensive results, users can run DENBTest first and then DEpermutePval, which generates results as follows.

```
> DEGres <- DENBTest(DEG)
```

- > DEGres <- DEpermutePval(DEGres, DEpermNBstat)
- > DEGres[1:6, c("NBstat", "pval", "padj", "perm.pval", "perm.padj")]

|                 | NBstat     | pval         | padj         | perm.pval | perm.padj |
|-----------------|------------|--------------|--------------|-----------|-----------|
| ENSG0000000003  | 0.5426504  | 3.956985e-01 | 5.408276e-01 | 0.55      | 1         |
| ENSG0000000005  | 0.2503510  | 3.300042e-01 | 4.943803e-01 | 0.30      | 1         |
| ENSG00000000419 | 0.0231052  | 9.244775e-01 | 9.839468e-01 | 0.95      | 1         |
| ENSG00000000457 | 14.3384053 | 9.960426e-05 | 2.589711e-03 | 0.00      | 0         |
| ENSG00000000460 | 1.4101270  | 1.370959e-01 | 2.970412e-01 | 0.40      | 1         |
| ENSG00000000938 | 2.1976989  | 7.309013e-07 | 4.434134e-05 | 0.00      | 0         |

# 4 Integrative GSEA runs

#### 4.1 DE/DS score integration

We have proposed two strategies for integrating normalized DE and DS scores (Wang and Cairns, 2013), one of which is the weighted summation of the two scores and the other is a rank-based strategy. The functions geneScore and genePermuteScore implement two methods for the weighted summation strategy: weighted linear combination and weighted quadratic combination. Scripts below show a linear combination of DE and DS scores with weight for DE equal to 0.3. Users should keep the weight for DE in geneScore and genePermuteScore the same, and the weight rangs from 0 (i.e., DS only) to 1 (i.e., DE only). Visualization of gene scores can be made by applying the plotGeneScore function.

```
> gene.score <- geneScore(DEscore, DSscore, method="linear", DEweight = 0.3)
> gene.score.perm <- genePermuteScore(DEscore.perm, DSscore.perm,
+ method="linear", DEweight=0.3)
> plotGeneScore(gene.score, gene.score.perm)
```

The plot generated by the plotGeneScore function (Fig. 1) can also be saved as a PDF file easily with the pdf argument of plotGeneScore.

The functions geneScore and genePermuteScore also implement one method for the rank-based integration strategy: using data-set-specific ranks. The plot for integrated gene scores is shown in Fig. 2.

#### **Gene List Overall Scoring Profile**

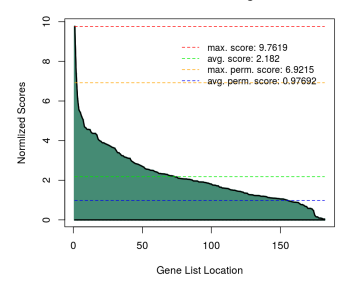


Figure 1: Gene scores resulted from linear combination. Scores are sorted from the largest to the smallest. Red, green, orange, blue dotted horizontal lines represent the maximum score, average score on the real data set, and the maximum score, average score on the permutation data sets.

```
> gene.score <- geneScore(DEscore, DSscore, method="rank", DEweight = 0.3)
> gene.score.perm <- genePermuteScore(DEscore.perm, DSscore.perm,
+ method="rank", DEweight=0.3)
> plotGeneScore(gene.score, gene.score.perm)
```

Rather than the above method to integrate scores with data-set-specific ranks, an alternative method is implemented with the rankCombine function, which takes only the ranks from the real data set for integrating DE and DS scores on both real and permutation data sets. This provides a method in a global manner. The plot of gene scores is shown in Fig. 3.

```
> combine <- rankCombine(DEscore, DSscore, DEscore.perm, DSscore.perm, DEweight=0.3)
> gene.score <- combine$geneScore
> gene.score.perm <- combine$genePermuteScore</pre>
```

> plotGeneScore(gene.score, gene.score.perm)

Basically the integrated gene scores are distributed similarly with the three integration methods at DE weight 0.3 (Figs. 1, 2, and 3); however, according to the analysis in Wang and Cairns (2013), SeqGSEA can detect slightly more significant gene sets with rank-based integration strategy than with linear combination.

#### 4.2 Initialization of SegGeneSet objects

To facilitate running gene set enrichment analysis, SeqGSEA implements a SeqGeneSet class. The SeqGeneSet class has several slots for accommodating a category of gene sets derived from any biological knowledge-based databases such as Kyoto Encyclopedia of Genes and Genomes (KEGG). However, we recommend to start with the formatted gene-set files from the well-maintained resource Molecular Signatures Database (MSigDB, http://www.broadinstitute.org/gsea/msigdb/index.jsp) (Subramanian et al., 2005). After downloading a gmt file from the above URL, users can use

#### **Gene List Overall Scoring Profile**

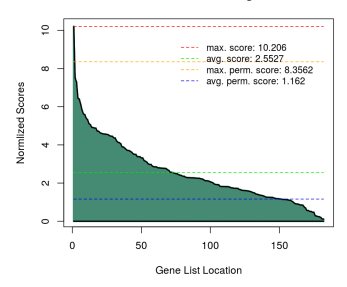


Figure 2: Gene scores resulted from rank-based combination with data-set-specific ranks. Scores are sorted from the largest to the smallest. Red, green, orange, blue dotted horizontal lines represent the maximum score, average score on the real data set, and the maximum score, average score on the permutation data sets.

#### **Gene List Overall Scoring Profile**

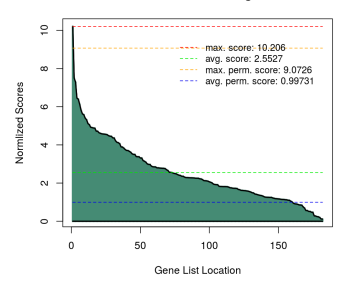


Figure 3: Gene scores resulted from rank-based combination with the same rank got from the real data set. Scores are sorted from the largest to the smallest. Red, green, orange, blue dotted horizontal lines represent the maximum score, average score on the real data set, and the maximum score, average score on the permutation data sets.

loadGenesets to initialize a SeqGeneSet object easily. Please note that with the current version of SeqGSEA, only gene sets with gene symbols are supported, though read count data's gene IDs can be either gene symbols or Ensembl Gene IDs.

Below is shown an example of the SeqGeneSet object, which contains information such as how many gene sets in this object and the names/sizes/descriptions of each gene set.

```
> data(GS_example, package="SeqGSEA")
> GS_example
SeqGeneSet object: gs_symb.txt
GeneSetSourceFile: /Library/Frameworks/R.framework/Versions/2.15/Resources/library/SeqGSEA/extdat
GeneSets: ERB2_UP.V1_DN
          AKT_UP_MTOR_DN.V1_UP
          KRAS.600.LUNG.BREAST_UP.V1_DN
 with the number of genes in respective sets: 6, 6, ..., 5
 brief descriptions:
          http://www.broadinstitute.org/gsea/msigdb/cards/ERB2_UP.V1_DN
          http://www.broadinstitute.org/gsea/msigdb/cards/AKT_UP_MTOR_DN.V1_UP
          http://www.broadinstitute.org/gsea/msigdb/cards/KRAS.600.LUNG.BREAST_UP.V1_DN
  # gene sets passed filter: 11 (#genes >= 5 AND <= 1000)</pre>
  # gene sets excluded: 178 (#genes < 5 OR > 1000)
ES scores: not computed
ES postions: not computed
Permutated ES scores: not performed
ES scores normalized: No
ES p-value: not computed
ES FWER: not computed
ES FDR: not computed
```

#### 4.3 running GSEA with integrated gene scores

With the initialized SeqGeneSet object and integrated gene scores as well as gene scores on the permutation data sets, the main GSEnrichAnlyze can be executed; and the topGeneSets allows users promptly access to the top significant gene sets.

```
> GS_example <- GSEnrichAnalyze(GS_example, gene.score, gene.score.perm)
> topGeneSets(GS_example, 5)
```

|    | GSName                        | ${\tt GSSize}$ | ES        | ES.pos | pval | FDR     | FWER |
|----|-------------------------------|----------------|-----------|--------|------|---------|------|
| 9  | TBK1.DF_UP                    | 5              | 1.8177717 | 17     | 0.05 | 0.00000 | 0.15 |
| 11 | KRAS.600.LUNG.BREAST_UP.V1_DN | 5              | 1.3045162 | 55     | 0.30 | 0.50000 | 0.95 |
| 5  | PKCA_DN.V1_DN                 | 5              | 1.4024471 | 74     | 0.30 | 0.66667 | 0.90 |
| 7  | BMI1_DN.V1_DN                 | 6              | 1.1377391 | 61     | 0.40 | 0.70000 | 1.00 |
| 6  | PTEN DN.V2 DN                 | 5              | 0.7987392 | 61     | 0.65 | 0.83333 | 1.00 |

The main GSEA includes several steps detailed in Wang and Cairns (2013) and its original paper Subramanian et al. (2005). In SeqGSEA, functions cales, cales.perm, normes and signifes are implemented to complete the analysis. Advanced users may set up customized pipelines with the functions above themselves.

#### Global Observed and Null Densities (Area Normalized)

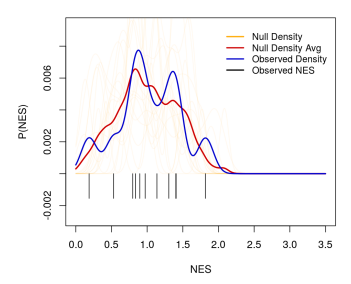


Figure 4: Distribution of normalized enrichment scores (NES) on the observed and permutation (null) data sets. Blue: observed NES density; Orange: each for NES density on one permutation data set; Red: the average density on all permutation data sets; Black: observed NES values.

### 4.4 SeqGSEA result displays

Several functions in SeqGSEA can be employed for visualization of gene set enrichment analysis running results. The plotES function is to plot the distribution of normalized enrichment scores (NES) of all gene sets in a SeqGeneSet object on the observed data set versus its empirical background provided by the NES on the permutation data sets (Fig. 4).

#### > plotES(GS\_example)

The plotSig function plots the distributions of permutation p-value, false discovery rate (FDR) and family-wise error rate (FWER) versus NES. The example plot is not shown in this vignette as the distributions can be far from the real ones due to the limited permutation times.

#### > plotSig(GS\_example)

The plotSigGS function is to plot detailed results of a particular gene set that has been analyzed. Information in the plot includes running enrichment scores, null NES on the permutation data sets. See Fig. 5 for an example.

> plotSigGeneSet(GS\_example, 9, gene.score) # 9th gene set is the most significant one.

Besides the functions to generate plots, the writeSigGeneSet function can write the detailed information of any analyzed gene sets, including NES, p-values, FDR, and the leading set (see the definition in Wang and Cairns (2013)). An example is shown below.

> writeSigGeneSet(GS\_example, 9, gene.score) # 9th gene set is the most significant one.

```
GSEA result for gene set No. 9:
genesetName gs_symb.txt:TBK1.DF_UP
```

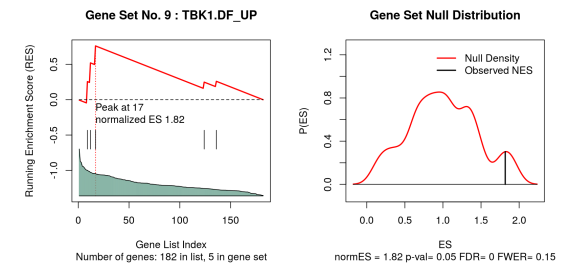


Figure 5: Left: gene locations of a particular gene set according to the gene score rank and running enrichment scores; Right: null NES distribution and the relative position of the observed NES.

genesetSize genesetDesc http://www.broadinstitute.org/gsea/msigdb/cards/TBK1.DF\_UP NES 1.8177717431103 Pos 17 pvalue 0.05 0 FDR **FWER** 0.15 Leading set: ENSG00000002919 5.58194069598269 ENSG00000005194 5.11651995163508 ENSG0000001167 4.87451465116361 Whole gene set: ENSG00000002919 5.58194069598269 ENSG0000005194 5.11651995163508 ENSG0000001167 4.87451465116361 ENSG0000005059 1.61250550921232 ENSG00000006576 1.36707987329728

The GSEAresultTable generates a summary table of the GSEA analysis, which can also be output with customized scripts. An example can be found in Section 6.

# 5 Running SeqGSEA with multiple cores

#### 5.1 R-parallel packages

There are many R packages for facilitating users in running R scripts in parallel, including *parallel*, *snowfall*, *multicore*, and many others. While experienced users may parallelize *SeqGSEA* runnings with the above packages themselves to reduce the running time, we provide with in the *SeqGSEA* 

package vignette an general way for users to parallelize their runnings utilizing the doParallel package (which depends on parallel).

First, we show a toy example for a basic idea how doParallel works. Basically, doParallel is a parallel backend for the foreach package using parallel, which provides a mechanism to execute foreach loops in parallel. With the foreach function in the foreach package, we can specify which foreach loops need to be parallelized using the %dopar% operator. However, without a registered parallel backend, the foreach loops will be executed sequentially even if the %dopar% operator is used. In those cases, the foreach package will issue a warning that it is running sequentially. Below are two running examples showing how the task is running sequentially and in parallel, respectively.

#### Run sequentially without parallel backend registered

```
> library(doParallel)
> a <- matrix(1:16, 4, 4)
> b \leftarrow t(a)
> foreach(b=iter(b, by='col'), .combine=cbind) %dopar%
    (a %*% b)
     [,1] [,2] [,3] [,4]
     276
[1,]
           304
                332
                      360
[2,]
     304
           336
                368
                      400
[3,]
      332
           368
                 404
                      440
[4,]
      360
           400
                440
                      480
```

Although the warning message didn't appear here, you would definitely see a warning message when you run the scripts above, like:

Warning message:

executing %dopar% sequentially: no parallel backend registered

#### Run in parallel with two cores

```
> library(doParallel)
> cl <- makeCluster(2) # specify 2 cores to be used in this computing
> registerDoParallel(cl)
> getDoParWorkers() # 2
[1] 2
> a <- matrix(1:16, 4, 4)
> b <- t(a)
> foreach(b=iter(b, by='col'), .combine=cbind) %dopar%
    (a \% * \% b)
     [,1] [,2] [,3] [,4]
[1,] 276
          304
                332
                     360
[2,]
      304
           336
                368
                     400
[3,]
      332
           368
                404
                     440
           400
[4,]
     360
                440
```

The parallel backend registration was done with registerdoParallel. For more details please refer to doParallel's vignette (http://cran.r-project.org/web/packages/doParallel/index.html).

#### 5.2 Parallelizing analysis on permutation data sets

In SeqGSEA, the loops for analyzing permutation data sets are implemented by foreach with %dopar% operator used. Those loops include DS, DE, and GSEA analyses, which are the most time consuming parts. Although there are three parts can take the advantage of parallel running, users only need to register parallel backend once at the beginning of all analyses. See an analysis example in the next section (Section 6).

# 6 Analysis examples

#### 6.1 Starting from your own RNA-Seq data

With this SeqGSEA package, we provide complementary Python scripts for counting reads on exons of each genes from SAM/BAM files: two scripts prepare\_exon\_annotation\_refseq.py and prepare\_exon\_annotation\_ensembl.py for preparing (sub-)exon annotation, and count\_in\_exons.py for counting reads. The scripts are based on the HTSeq Python package (http://www-huber.embl.de/users/anders/HTSeq/). Please install it before using the Python scripts provided. The scripts can be found in the directory given by the following command.

```
> system.file("extscripts", package="SeqGSEA", mustWork=TRUE)
```

#### [1] "/tmp/RtmpFPsHvt/Rinst5aaa652a0aa4/SeqGSEA/extscripts"

Simply by typing "python" + the file name of echo script in your shell console, the help documentation will be on your screen.

Other than the Python scripts provided, users who prefer playing with R/Bioconductor packages can also use easyRNASeq in easyRNASeq, summarizeOverlaps in GenomicRanges, and featureCounts in Rsubread to count reads that mapped to each exon. Please refer to respective packages for detailed usage.

For users who are not familiar with RNA-Seq data processing, the upstream steps of counting reads are (1) data preprocessing, including adapter removal, low-quality read filtering, data quality-control analysis, and (2) read mapping. R/Bioconductor users can apply Rsubread to map reads based on a seed-and-vote approach, as well as a few QC analysis. Users familiar with command-line can choose from a wide range of tools, such as already widely used ones including TopHat (http://tophat.cbcb.umd.edu), START (http://code.google.com/p/rna-star), and etc..

#### 6.2 Exemplified pipeline for integrating DE and DS

Below is shown a typical SeqGSEA running example with the data enclosed with the SeqGSEA package, which are a part of the prostate cancer data set (Kannan et al., 2011). We divide the process into five steps for a complete SeqGSEA run.

Step 0: Initialization. (Users should change values in this part accordingly.)

```
> # output file prefix
> output.prefix <- "SeqGSEA.test"</pre>
> # setup parallel backend
> library(doParallel)
> cl <- makeCluster(2) # specify 2 cores to be used in computing
> registerDoParallel(cl) # parallel backend registration
> # setup permutation times
> perm.times <- 20 # change the number to >= 1000 in your analysis
Step 1: DS analysis
> # load exon read count data
> RCS <- loadExonCountData(case.files, control.files)
> # remove genes with low expression
> RCS <- exonTestability(RCS, cutoff=5)</pre>
> geneTestable <- geneTestability(RCS)
> RCS <- subsetByGenes(RCS, unique(geneID(RCS))[ geneTestable ])</pre>
> # get gene IDs, which will be used in initialization of gene set
> geneIDs <- unique(geneID(RCS))</pre>
> # calculate DS NB statistics
> RCS <- estiExonNBstat(RCS)
> RCS <- estiGeneNBstat(RCS)
> # calculate DS NB statistics on the permutation data sets
> permuteMat <- genpermuteMat(RCS, times=perm.times)</pre>
> RCS <- DSpermute4GSEA(RCS, permuteMat)
Step 2: DE analysis
> # get gene read counts
> geneCounts <- getGeneCount(RCS)
> # calculate DE NB statistics
> label <- label(RCS)</pre>
> DEG <-runDESeq(geneCounts, label)
> DEGres <- DENBStat4GSEA(DEG)
> # calculate DE NB statistics on the permutation data sets
> DEpermNBstat <- DENBStatPermut4GSEA(DEG, permuteMat) # permutation
Step 3: score integration
> # DE score normalization
> DEscore.normFac <- normFactor(DEpermNBstat)</pre>
> DEscore <- scoreNormalization(DEGres$NBstat, DEscore.normFac)
> DEscore.perm <- scoreNormalization(DEpermNBstat, DEscore.normFac)
> # DS score normalization
> DSscore.normFac <- normFactor(RCS@permute_NBstat_gene)</pre>
> DSscore <- scoreNormalization(RCS@featureData_gene$NBstat, DSscore.normFac)
> DSscore.perm <- scoreNormalization(RCS@permute_NBstat_gene, DSscore.normFac)
> # score integration
> gene.score <- geneScore(DEscore, DSscore, DEweight=0.5)
> gene.score.perm <- genePermuteScore(DEscore.perm, DSscore.perm, DEweight=0.5)
> # visilization of scores
> # NOT run in the example; users to uncomment the following 6 lines to run
> #plotGeneScore(DEscore, DEscore.perm, pdf=paste(output.prefix,".DEScore.pdf",sep=""),
                 main="Expression")
> #plotGeneScore(DSscore, DSscore.perm, pdf=paste(output.prefix,".DSScore.pdf",sep=""),
```

```
> #
                 main="Splicing")
> #plotGeneScore(gene.score, gene.score.perm,
                 pdf=paste(output.prefix,".GeneScore.pdf",sep=""))
Step 4: main GSEA
> # load gene set data
> gene.set <- loadGenesets(geneset.file, geneIDs, geneID.type="ensembl",
                           genesetsize.min = 5, genesetsize.max = 1000)
> # enrichment analysis
> gene.set <- GSEnrichAnalyze(gene.set, gene.score, gene.score.perm, weighted.type=1)
> # format enrichment analysis results
> GSEAres <- GSEAresultTable(gene.set, TRUE)
> # output results
> # NOT run in the example; users to uncomment the following 4 lines to run
> #write.table(GSEAres, paste(output.prefix, ".GSEA.result.txt", sep=""),
               quote=FALSE, sep="\t", row.names=FALSE)
> #plotES(gene.set, pdf=paste(output.prefix,".GSEA.ES.pdf",sep=""))
> #plotSig(gene.set, pdf=paste(output.prefix,".GSEA.FDR.pdf",sep=""))
```

For gene sets used in Step 4, while we recommend users directly download and use those already well-formatted gene sets from MSigDB (http://www.broadinstitute.org/gsea/msigdb/index.jsp), users can also feed whatever gene sets to SeqGSEA as long as they are in the GMT format. Please refer to the following URL for details: http://www.broadinstitute.org/cancer/software/gsea/wiki/index.php/Data\_formats.

## 6.3 Exemplified pipeline for DE-only analysis

For the demanding of DE-only analysis, such as for organisms without much alternative splicing annotated, here we show an exemplified pipeline for such analysis. It includes 4 steps as follows.

**Step 0:** Initialization. (Users should change values in this part accordingly.)

```
> rm(list=ls())
> # input count data files
> data.dir <- system.file("extdata", package="SeqGSEA", mustWork=TRUE)</pre>
> count.file <- paste(data.dir, "geneCounts.txt", sep="/")</pre>
> # gene set file
> geneset.file <- system.file("extdata", "gs_symb.txt",
                              package="SeqGSEA", mustWork=TRUE)
> # output file prefix
> output.prefix <- "SeqGSEA.test"
> # setup parallel backend
> library(doParallel)
> cl <- makeCluster(2) # specify 2 cores to be used in computing
> registerDoParallel(cl) # parallel backend registration
> # setup permutation times
> perm.times <- 20 # change the number to >= 1000 in your analysis
Step 1: DE analysis
> # load gene read count data
> geneCounts <- read.table(count.file)</pre>
> # speficify the labels of each sample
> label <- as.factor(c(rep(1,10), rep(0,10)))
```

```
> # calculate DE NB statistics
> DEG <-runDESeq(geneCounts, label)
> DEGres <- DENBStat4GSEA(DEG)
> # calculate DE NB statistics on the permutation data sets
> permuteMat <- genpermuteMat(label, times=perm.times)</pre>
> DEpermNBstat <- DENBStatPermut4GSEA(DEG, permuteMat) # permutation
Step 2: score normalization
> # DE score normalization
> DEscore.normFac <- normFactor(DEpermNBstat)</pre>
> DEscore <- scoreNormalization(DEGres$NBstat, DEscore.normFac)
> DEscore.perm <- scoreNormalization(DEpermNBstat, DEscore.normFac)
> # score integration - DSscore can be null
> gene.score <- geneScore(DEscore, DEweight=1)</pre>
> gene.score.perm <- genePermuteScore(DEscore.perm, DEweight=1) # visilization of scores
> # NOT run in the example; users to uncomment the following 6 lines to run
> #plotGeneScore(DEscore, DEscore.perm, pdf=paste(output.prefix,".DEScore.pdf",sep=""),
                 main="Expression")
> #plotGeneScore(gene.score, gene.score.perm,
                 pdf=paste(output.prefix,".GeneScore.pdf",sep=""))
Step 3: main GSEA
> # load gene set data
> geneIDs <- rownames(geneCounts)</pre>
> gene.set <- loadGenesets(geneset.file, geneIDs, geneID.type="ensembl",
                           genesetsize.min = 5, genesetsize.max = 1000)
> # enrichment analysis
> gene.set <- GSEnrichAnalyze(gene.set, gene.score, gene.score.perm, weighted.type=1)
> # format enrichment analysis results
> GSEAres <- GSEAresultTable(gene.set, TRUE)
> # output results
> # NOT run in the example; users to uncomment the following 4 lines to run
> #write.table(GSEAres, paste(output.prefix,".GSEA.result.txt",sep=""),
> #
               quote=FALSE, sep="\t", row.names=FALSE)
> #plotES(gene.set, pdf=paste(output.prefix,".GSEA.ES.pdf",sep=""))
> #plotSig(gene.set, pdf=paste(output.prefix,".GSEA.FDR.pdf",sep=""))
```

#### 6.4 One-step SeqGSEA analysis

While users can choose to run SeqGSEA step by step in a well-controlled manner (see above), the one-step SeqGSEA analysis with an all-in runSeqGSEA function enables users to run SeqGSEA in the easiest way. With the runSeqGSEA function, users can also test multiple weights for integrating DE and DS scores. DE-only analysis starting with exon read counts is also supported in the all-in function.

Follow the example below to start your first SeqGSEA analysis now!

```
> ### Initialization ###
> # input file location and pattern
> data.dir <- system.file("extdata", package="SeqGSEA", mustWork=TRUE)
> case.pattern <- "^SC" # file name starting with "SC"
> ctrl.pattern <- "^SN" # file name starting with "SN"
> # gene set file and type
> geneset.file <- system.file("extdata", "gs_symb.txt",</pre>
```

```
package="SeqGSEA", mustWork=TRUE)
> geneID.type <- "ensembl"</pre>
> # output file prefix
> output.prefix <- "SeqGSEA.example"
> # analysis parameters
> nCores <- 8
> perm.times <- 1000 # >= 1000 recommended
> DEonly <- FALSE
> DEweight \langle -c(0.2, 0.5, 0.8) \rangle # a vector for different weights
> integrationMethod <- "linear"</pre>
> ### one step SeqGSEA running ###
> # NOT run in the example; uncomment the following 4 lines to run
> # CAUTION: running the following lines will generate lots of files in your working dir
> #runSeqGSEA(data.dir=data.dir, case.pattern=case.pattern, ctrl.pattern=ctrl.pattern,
              geneset.file=geneset.file, geneID.type=geneID.type, output.prefix=output.prefix,
              {\tt nCores=nCores}, \ {\tt perm.times=perm.times}, \ {\tt integrationMethod=integrationMethod},
> #
> #
              DEonly=DEonly, DEweight=DEweight)
    Session information
> sessionInfo()
R version 3.4.2 (2017-09-28)
Platform: x86_64-pc-linux-gnu (64-bit)
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
locale:
 [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
                                LC_COLLATE=C
 [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 stats
                        graphics grDevices utils
                                                       datasets methods
[8] base
other attached packages:
[1] SeqGSEA_1.18.0
                        DESeq_1.30.0
                                             lattice_0.20-35
[4] locfit_1.5-9.1
                        doParallel_1.0.11
                                            iterators_1.0.8
[7] foreach_1.4.3
                        Biobase_2.38.0
                                            BiocGenerics_0.24.0
loaded via a namespace (and not attached):
                                                RColorBrewer_1.1-2
 [1] Rcpp_0.12.13
                         compiler_3.4.2
 [4] prettyunits_1.0.2
                          progress_1.1.2
                                                bitops_1.0-6
 [7] tools_3.4.2
                          biomaRt_2.34.0
                                                digest_0.6.12
[10] bit_1.1-12
                                                RSQLite_2.0
                          annotate_1.56.0
```

```
[13] memoise_1.1.0
                          tibble_1.3.4
                                                rlang_0.1.2
[16] Matrix_1.2-11
                          DBI_0.7
                                                stringr_1.2.0
[19] genefilter_1.60.0
                          S4Vectors_0.16.0
                                                IRanges_2.12.0
[22] stats4_3.4.2
                          bit64_0.9-7
                                                grid_3.4.2
[25] R6_2.2.2
                          AnnotationDbi_1.40.0 XML_3.98-1.9
[28] survival_2.41-3
                          magrittr_1.5
                                                geneplotter_1.56.0
[31] blob_1.1.0
                          codetools_0.2-15
                                                splines_3.4.2
[34] assertthat_0.2.0
                          xtable_1.8-2
                                                stringi_1.1.5
[37] RCurl_1.95-4.8
```

# Cleanup

This is a cleanup step for the vignette on Windows; typically not needed for users.

```
> allCon <- showConnections()
> socketCon <- as.integer(rownames(allCon)[allCon[, "class"] == "sockconn"])
> sapply(socketCon, function(ii) close.connection(getConnection(ii)))
```

#### References

- Anders, S. and Huber, W. (2010). Differential expression analysis for sequence count data. *Genome Biology*, 11:R106.
- Benjamini, Y. and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. Ser. B, 57(1):289–300.
- Kannan, K., Wang, L., Wang, J., Ittmann, M. M., Li, W., and Yen, L. (2011). Recurrent chimeric rnas enriched in human prostate cancer identified by deep sequencing. *Proc Natl Acad Sci U S A*, 108(22):9172–7.
- Subramanian, A., Tamayo, P., Mootha, V. K., Mukherjee, S., Ebert, B. L., Gillette, M. A., Paulovich, A., Pomeroy, S. L., Golub, T. R., Lander, E. S., and Mesirov, J. P. (2005). Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. *Proc Natl Acad Sci U S A*, 102(43):15545–50.
- Wang, W., Qin, Z., Feng, Z., Wang, X., and Zhang, X. (2013). Identifying differentially spliced genes from two groups of rna-seq samples. *Gene*, 518(1):164–170.
- Wang, X. and Cairns, M. (2013). Gene set enrichment analysis of RNA-Seq data: integrating differential expression and splicing. *BMC Bioinformatics*, 14(Suppl 5):S16.
- Wang, X. and Cairns, M. (2014). SeqGSEA: a Bioconductor package for gene set enrichment analysis of RNA-Seq data integrating differential expression and splicing. *Bioinformatics*, 30(12):1777–9.