

# Package ‘RTopper’

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**Type** Package

**Title** This package is designed to perform Gene Set Analysis across multiple genomic platforms

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**Depends** R (>= 2.12.0), Biobase

**Imports** limma, multtest

**Suggests** org.Hs.eg.db, KEGGREST, GO.db

**Description** the RTopper package is designed to perform and integrate gene set enrichment results across multiple genomic platforms.

**License** GPL (>= 3) + file LICENSE

**biocViews** Microarray

**LazyLoad** yes

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|                 |                                                                              |
|-----------------|------------------------------------------------------------------------------|
| RTopper-package | <i>A package to perform run Gene Set Enrichment across genomic platforms</i> |
|-----------------|------------------------------------------------------------------------------|

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**Description**

Gene sets analysis considers whether genes sharing a biological property also behave in a related way in experimental data. This technique is commonly used in high throughput genomic analyses to assist results interpretation, and has been successfully applied in cancer genome projects for integrating information from multiple genome-wide assays. The RTopper package uses gene sets analysis to overcome the diversity of genomic data providing the statistical framework for integration across data types.

**Details**

Package: RTopper  
Type: Package  
Version: 0.1  
Date: 2010-11-12  
License: GPL version 3 or newer

**RTopper package features**

- Rtopper enables two gene set-based data integration approaches:
- 1 Integration+GSA: computing integrated gene-to-phenotype association scores, followed by conventional gene sets analysis;
  - 2 GSA+Integration: computing consensus significance score after all data types are analyzed individually;
  - 3 Use of alternative enrichment test: RTopper uses the Wilcoxon rank-sum test for enrichment testing, however alternative tests can be defined and used;
  - 4 Multiple testing correction: RTopper enables adjustment of p-values obtained from enrichment analysis;

**Author(s)**

Luigi Marchionni <marchion@jhu.edu>

**References**

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani. "Integrating diverse genomic data using gene sets." Manuscript submitted.

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|               |                                    |
|---------------|------------------------------------|
| adjustPvalGSE | <i>Multiple testing correction</i> |
|---------------|------------------------------------|

---

**Description**

This function is an interface to the `mt.rawp2adjp` function contained in the `multtest` package. `adjustPvalGSE` works on outputs from `runBatchGSE` and `combineGSE` returning adjusted p-values

**Usage**

```
adjustPvalGSE(gseOut, proc = "BH", alpha = 0.05 , na.rm = FALSE)
```

**Arguments**

|                     |                                                                                                                                                                                                                                                                                                                  |
|---------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>gseOut</code> | list of lists, either the output from <code>runBatchGSE</code> , or the output from <code>combineGSE</code>                                                                                                                                                                                                      |
| <code>proc</code>   | character, the method to be used for p-values adjusting. This parameter will be passed to the <code>mt.rawp2adjp</code> function from the <code>multtest</code> package. The available options include: "Bonferroni", "Holm", "Hochberg", "SidakSS", "code"SidakSD", "BH" (the default), "BY", "ABH", and "TSBH" |
| <code>alpha</code>  | numeric, the nominal type I error rate                                                                                                                                                                                                                                                                           |
| <code>na.rm</code>  | logical, the option for handling NA values in the list of raw p-values                                                                                                                                                                                                                                           |

**Details**

The `adjustPvalGSE` function performs p-value adjusting for multiple testing correction on the list of lists resulting from enrichment analysis obtained using the `runBatchGSE` and `combineGSE` functions. This functions is based on the `mt.rawp2adjp` function contained in the `multtest` package.

**Value**

For each vector of p-value contained in the `gseOut` input object a data.frame is returned, containing original p-value and corrected p-values

**Author(s)**

Luigi Marchionni <marchion@jh.edu>

## References

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani. "Integrating diverse genomic data using gene sets." Manuscript submitted.

## Examples

```
###load gse analysis results for separate gene-to-phenotype score
data(gseResultsSep)

###adjust for multiple testing using the Benjamini and Hochberg method
gseABS.int.BH <- adjustPvalGSE(gseResultsSep)

###adjust for multiple testing using the Holm method
gseABS.int.holm <- adjustPvalGSE(gseResultsSep, proc = "Holm")
```

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|            |                             |
|------------|-----------------------------|
| combineGSE | <i>Combines GSE results</i> |
|------------|-----------------------------|

---

## Description

Combines GSE results obtained from a separate set of gene-to-phenotypes scores

## Usage

```
combineGSE(gseOut, method)
```

## Arguments

|        |                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
|--------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| gseOut | a list of lists containing the enrichment results to be combined. This is usually the the output of <a href="#">runBatchGSE</a> obtained from a set of distinct genes-to-phenotype scores (usually one per genomic platform)                                                                                                                                                                                                                                |
| method | character, this argument specifies the method used to combine the enrichment results obtained from distinct genes-to-phenotype scores (usually one per genomic platform). Available options are the computation of the geometric or arithmetic means, the use of the median, the selection of the minimum or the maximum enrichment score, and the random selection of a score (respectively "geometricMean", "mean", "median", "min", "max", and "random") |

## Details

This function summarize enrichment results obtained from distinct gene-to-phenotypes scores (usually one per genomic platform) by one of several alternative methods.

## Value

The output is a list of lists containing integrated enrichment results for all FGS collections

**Author(s)**

Luigi Marchionni <marchion@jhu.edu>

**References**

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani. "Integrating diverse genomic data using gene sets." Manuscript submitted.

**Examples**

```
###load gse analysis results for separate gene-to-phenotype score
data(gseResultsSep)

###combine enrichment score results using geometric mean
gseABS.sep.geoMean <- combineGSE(gseResultsSep, method="geometricMean")

###combine enrichment score results using maximum value
gseABS.sep.max <- combineGSE(gseResultsSep, method="max")
```

---

computeDrStat

*Computes gene-to-phenotype associations scores*

---

**Description**

computeDrStat Computes gene-to-phenotype associations scores, using as input the output from convertToDr.

**Usage**

```
computeDrStat(data, columns = c(1:(ncol(data)-1)), method = "dev", integrate = TRUE)
```

**Arguments**

|           |                                                                                                                                                             |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data      | a list of data.frames containing genomic measurements. Each element of dataIntersection must account for the same set of patients(columns) and genes (rows) |
| columns   | a data.frame indicating patients' phenotypic class                                                                                                          |
| method    | character, the number of genomic platforms                                                                                                                  |
| integrate | logical, wheter to integrate the gene-to-phenotype scores across platform or return separates scores for each platform                                      |

## Details

This function allows computing gene-to-phenotype association scores, using as input the gene-centered list produced by `computeDr`. The `computeDrStat` function works separately on each gene-centered data.frame created by the `convertToDr` function, assuming that the phenotype information is stored in the last column named "response". It is possible computing both separate association scores for each platform, as well as an integrated score, as specified by the `integrate` arguments. There are currently three methods available for obtaining the scores (see Tyekucheva et al, manuscript under review), as specified by the `methods` argument:

"dev": this approach computes the score as the difference of deviances;

"aic": this approach computes the score as the Akaike information criterion for model selection;

"bic": this approach computes the score as the penalized likelihood ratio;

## Value

A list of named vectors containing separate or integrated gene-to-phenotype association scores.

## Author(s)

Luigi Marchionni <marchion@jhu.edu>

## References

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani. "Integrating diverse genomic data using gene sets." Manuscript submitted.

## Examples

```
###load data
data(exampleData)

###convert
dataDr <- convertToDr(dat, pheno, 4)

###compute the integrated score
bicStatIntegrated <- computeDrStat(dataDr, columns = c(1:4), method="bic", integrate = TRUE)

###compute separate scores for each genomic platform
bicStatSeparate <- computeDrStat(dataDr, columns = c(1:4), method="bic", integrate = FALSE)
```

---

|             |                                                                                        |
|-------------|----------------------------------------------------------------------------------------|
| convertToDr | <i>Converts genomic data to a list suitable for computing gene-to-phenotype scores</i> |
|-------------|----------------------------------------------------------------------------------------|

---

## Description

`convertToDr` converts genomic data into a list further used for computing gene-to-phenotype association scores.

**Usage**

```
convertToDr(dataIntersection, response, nPlatforms = length(data))
```

**Arguments**

|                  |                                                                                                                                                             |
|------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------|
| dataIntersection | a list of data.frames containing genomic measurements. Each element of dataIntersection must account for the same set of patients(columns) and genes (rows) |
| response         | a data.frame indicating patients' phenotypic class                                                                                                          |
| nPlatforms       | numeric, the number of genomic platforms                                                                                                                    |

**Details**

This function converts a list of data.frames containing distinct genomic measurements performed on the same patients into a gene-centered used in further analyses for computing gene-to-phenotype scores. Data.frame in the input list (dataIntersection) must have the same dimensions, with columns being patients, and rows being genes. Column names identify the patients, while rownames identify the genes. The argument response is used to pass phenotypic information about samples to be analyzed. This is a simple two columns data.frame in which the first column corresponds to patients identifiers, and the second column to the phenotypic response encoded as binary class (using the integers 0 and 1). The nPlatforms argument specifies the number of platforms that will be analyzed.

**Value**

A list of data.frames, one for each analyzed gene, summarizing all genomic measurements and phenotypic information across patients and platforms.

**Author(s)**

Luigi Marchionni <marchion@jhu.edu>

**References**

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani "Integrating diverse genomic data using gene sets." Manuscript submitted.

**Examples**

```
###load data
data(exampleData)

###convert
dataDr <- convertToDr(dat, pheno, 4)
```

---

`dat`*A test dataset for the RTopper package*

---

## Description

A small subset of pre-processed Glioblasoma Multiforme (GBM) genomic data from The Cancer Genome Atlas (TCGA) project, encompassing Differential Gene Expression (DGE), and Copy Number Variation (CNV). Can be used as input to [convertToDr](#).

## Usage

```
data(exampleData)
```

## Format

This object is a list of 4 `data.frames` containing genomic measurements obtained across distinct genomic scopes (copy number variation and gene expression), platforms (Affymetrix and Agiles), and laboratories. In particular each `data.frame` consist of 500 gene measurements (by rows), for 95 distinct patients (by columns) from the following 4 distinct platforms:

"`dat.affy`": DGE obtained using Affymetrix microarrays;

"`dat.agilent`": DGE obtained using Agilent microarrays;

"`dat.cnvHarvard`": CNV data obtained at Harvard;

"`dat.cnvMskcc`": CNV data obtained at Memorial Sloan Ketterng Cancer Center;

## Source

The Cancer Genome Atlas (TCGA) project <http://cancergenome.nih.gov/>

## References

The Cancer Genome Atlas (TCGA) Research Network. "Comprehensive genomic characterization defines human glioblastoma genes and core pathways". *Nature*, 2008, October 23; 455(7216): 1061-1068

## Examples

```
data(exampleData)
ls()
class(dat)
names(dat)
sapply(dat, class)
sapply(dat, dim)
```

---

|         |                                                                                                  |
|---------|--------------------------------------------------------------------------------------------------|
| fgsList | <i>A list of Functional Gene Set (FGS) to be used to run the examples in the RTopper package</i> |
|---------|--------------------------------------------------------------------------------------------------|

---

### Description

A list containing distinct types of FGS (i.e. Gene Ontology, KEGG pathways). Each FGS is type is a list of named character vectors, one for each FGS, containing the gene identifiers. Vectors names describe the FGS. Can be used as input to [runBatchGSE](#).

### Usage

```
data(fgsList)
```

### Format

This object is a list of length two:

"go": this is a list of 5 character vectors, corresponding to 5 distinct Gene Ontology (GO) terms. Genes annotated to each GO term are identified by their gene symbol;

"kegg": this is a list of 5 character vectors, corresponding to 5 distinct KEGG pathways. Genes annotated to each KEGG pathway are identified by their gene symbol;

### Source

The FGS were obtained from the `org.Hs.eg.db` package, (use the [org.Hs.eg](#) function to see the content); These FGS were annotated using data from `GO.db` and `KEGGREST` packages (use the [GO](#) and [keggGet](#) functions to see the content).

### Examples

```
data(fgsList)
class(fgsList)
names(fgsList)
str(fgsList)
```

---

|               |                                                                                                               |
|---------------|---------------------------------------------------------------------------------------------------------------|
| gseResultsSep | <i>A list of separated gene set enrichment p-values to be used to run the examples in the RTopper package</i> |
|---------------|---------------------------------------------------------------------------------------------------------------|

---

### Description

A list containing distinct named numeric vectors corresponding to the gene set enrichment p-value separately computed with [runBatchGSE](#) for each distinct data set. Note that for each data set there are two set of p-values, one for GO and one for KEGG. These separate p-values can be combined across data sets by the [combineGSE](#) function. Can be used as input to [adjustPvalGSE](#).

**Usage**

```
data(gseResultsSep)
```

**Format**

This object is a list of length four:

"dat.affy": a list of length two: "go" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 GO terms on Affymetrix gene expression data; "keg" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 KEGG pathways on Affymetrix gene expression data;

"dat.agilent": a list of length two: "go" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 GO terms on Agilent gene expression data; "keg" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 KEGG pathways on Agilent gene expression data;

"dat.cnvHarvard": a list of length two: "go" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 GO terms on Harvard CNV data; "keg" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 KEGG pathways on Harvard CNV data;

"dat.cnvMskcc": a list of length two: "go" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 GO terms on MSKCC CNV data; "keg" is a numeric vector of length 5, containing the p-values resulting from gene set enrichment analysis of 5 KEGG pathways on MSKCC CNV data;

**Source**

Computed using the [runBatchGSE](#) function from the TCGA data contained in [sepScores](#) and [fgsList](#).

**Examples**

```
data(gseResultsSep)
class(gseResultsSep)
names(gseResultsSep)
str(gseResultsSep)
```

---

intScores

*A list of genomic scores integrated across distinct data sets to be used to run the examples in the RTopper package*

---

**Description**

A list containing a named numeric vector corresponding to the genomic score resulting from the integration across distinct data set. These integrated gene-to-phenotype scores are computed by the [computeDrStat](#) function. Can be used as input to [runBatchGSE](#).

**Usage**

```
data(intScores)
```

**Format**

This object is a list of length one:

"integrated": a numeric vector of length 500, corresponding to the integrated phenotype association scores computed for each of the 500 genes used in the examples;

**Source**

Computed using the [computeDrStat](#) function from the TCGA data contained in [dat](#) and [pheno](#).

**Examples**

```
data(intScores)
class(intScores)
names(intScores)
str(intScores)
```

---

pheno

*A test dataset for the RTopper package*

---

**Description**

A `data.frame` with 2 columns containing the phenotypic class indicator for the 95 patients analyzed and used in the examples. Can be used as input to [convertToDr](#).

**Usage**

```
data(exampleData)
```

**Format**

This object is a `data.frame` with two columns:

"Sample": the first column contains the patients identifiers;

"Class": the second columns contain a numeric indicator (0 or 1) corresponding to the phenotypic class of each patient;

**Source**

The Cancer Genome Atlas (TCGA) project <http://cancergenome.nih.gov/>

**References**

The Cancer Genome Atlas (TCGA) Research Network. "Comprehensive genomic characterization defines human glioblastoma genes and core pathways". *Nature*, 2008, October 23; 455(7216): 1061-1068

## Examples

```
data(exampleData)
class(pheno)
colnames(pheno)
str(pheno)
```

---

|             |                                                                                |
|-------------|--------------------------------------------------------------------------------|
| runBatchGSE | <i>To perform GSE analysis over multiple experiments and functional themes</i> |
|-------------|--------------------------------------------------------------------------------|

---

## Description

The runBatchGSE function enables performing Gene Set Enrichment analysis over multiple ranking statistics and multiple lists of gene sets. By default this function is an interface to the [geneSetTest](#) in the limma package, and most of the arguments passed to runBatchGSE are indeed passed to such lower level function. As an alternative the user can also define and pass to runBatchGSE a custom function, defining the ranking statistics and the gene set membership in the same way done for [geneSetTest](#) (see Details below).

## Usage

```
runBatchGSE(dataList, fgsList, ...)
```

## Arguments

|          |                                                                                                                                                             |
|----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------|
| dataList | a list containing the gene-to-phenotype scores to be used as ranking statistics in the GSE analysis. This list is usually produced by running computeDrStat |
| fgsList  | a list of FGS collection, in which each element is a list of character vectors, one for each gene set                                                       |
| ...      | additional arguments to be passed to lower level functions (see details below)                                                                              |

## Details

This function performs enrichment analysis for all the gene-to-phenotype scores (argument dataList) passed to it over a list of Functional Gene Set (FGS) (argument fgsList), returning a p-value for each FGS. Additional arguments can be passed to this function to modify the way the enrichment test is performed, as follows:

absolute logical, this specifies whether the absolute values of the ranking statistics should be used in the test (the default being TRUE)

gseFunc a function to perform GSE analysis. If not specified the default is the [geneSetTest](#) function from the limma package. If a function is specified by the user, the membership of the analyzed genes to a FGS, and the ranking statistics must be defined in the same way this is done for [geneSetTest](#), and the new function must return an integer (usually a p-value) (see the help for [geneSetTest](#))

The following main arguments are used by [geneSetTest](#):

type character, specifies the type of statistics used to rank the genes by geneSetTest: 'f' for F-like statistics (default), 't' for t-like statistics, or 'auto' for an educated guess

alternative character, defines the alternative with the following possible options: 'mixed' (default), 'either', 'up' or 'down', 'two.sided', 'greater', or 'less'

ranks.only logical, if TRUE (default) only ranks will be used by geneSetTest

nsim numeric, the number of randomly selected sets of genes to be used in simulations to compute the p-value

## Value

The output is a list of lists containing the set of enrichment results for all gene-to-phenotype scores and FGS collections used as input.

## Author(s)

Luigi Marchionni <marchion@jhu.edu>

## References

Svitlana Tyekucheva, Luigi Marchionni, Rachel Karchin, and Giovanni Parmigiani. "Integrating diverse genomic data using gene sets." Manuscript submitted.

## Examples

```
###require limma to run the example
require(limma)

###load integrated gene-to-phenotype scores
data(intScores)

###load separate gene-to-phenotype scores
data(sepScores)

###load list of functional gene sets
data(fgsList)

###run GSE analysis in batch with default parameters
gseABS.int <- runBatchGSE(dataList=intScores, fgsList=fgsList)

###run GSE analysis in batch with alternative parameters
gseABS.sep <- runBatchGSE(dataList=sepScores, fgsList=fgsList,
  absolute=FALSE, type="t", alternative="up")

###run GSE analysis in batch passing an enrichment function
gseUP.int.2 <- runBatchGSE(dataList=intScores, fgsList=fgsList,
  absolute=FALSE, gseFunc=wilcoxGST, alternative="up")

###define and use a new enrichment function
gseFunc <- function (selected, statistics, threshold) {
  diffExpGenes <- statistics > threshold
  tab <- table(diffExpGenes, selected)
```

```

pVal <- fisher.test(tab)[["p.value"]]
}
gseUP.sep.2 <- runBatchGSE(dataList=sepScores, fgsList=fgsList,
  absolute=FALSE, gseFunc=gseFunc, threshold=7.5)

```

---

|           |                                                                                                                                                                         |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| sepScores | <i>A list of separate gene-to-phenotype association scores, obtained independently for each distinct data set to be used to run the examples in the RTopper package</i> |
|-----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

A list containing distinct named numeric vectors corresponding to the gene-to-phenotype association scores resulting from the separate analysis of each data set. These separate gene-to-phenotype scores are computed by [computeDrStat](#) function. Can be used as input to [runBatchGSE](#).

### Usage

```
data(sepScores)
```

### Format

This object is a list of length four, one element for data set:

"dat.affy": a numeric vector of length 500, corresponding to the separate phenotype association scores computed for Affymetrix gene expression data;

"dat.agilent": a numeric vector of length 500, corresponding to the separate phenotype association scores computed for Agilent gene expression data;

"dat.cnvHarvard": a numeric vector of length 500, corresponding to the separate phenotype association scores computed for Harvard CNV data;

"dat.cnvMskcc": a numeric vector of length 500, corresponding to the separate phenotype association scores computed for the MSKCC CNV data;

### Source

Computed using the [computeDrStat](#) function from the TCGA data contained in [data](#).

### Examples

```

data(sepScores)
class(sepScores)
names(sepScores)
str(sepScores)

```

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