# Package 'ROTS'

October 12, 2016

Type Package
Title Reproducibility-Optimized Test Statistic
Version 1.0.0
Author Fatemeh Seyednasrollah, Tomi Suomi, Laura L. Elo
Maintainer Fatemeh Seyednasrollah <fatsey@utu.fi></fatsey@utu.fi>
<b>Description</b> Calculates the Reproducibility-Optimized Test Statistic (ROTS) for differential testing in omics data.
<b>biocViews</b> Software, GeneExpression, DifferentialExpression, Microarray, RNASeq, Proteomics
License GPL (>= 2)
LazyData yes
LinkingTo Rcpp
Imports Rcpp, stats, Biobase, methods
<b>Depends</b> R (>= 3.3)
NeedsCompilation yes

Index

### R topics documented:

affySpikeIn	2
plot.ROTS	2
ROTS	3
summary.ROTS	5
upsSpikeIn	6
	7

1

```
affySpikeIn
```

#### Description

The Affymetrix HG-U95Av2 spike-in data contains two groups of 12 samples. In this carefully controlled experiment, it is known that 14 genes are differentially expressed between the two sample groups, while the rest of the genes are equally expressed. In order to reduce the running time of the vignette example, this package uses a subset of 1000 genes of the original dataset including all the spiked genes with 5 samples from each group. The preprocessed RMA-normalized data were obtained from the Bioconductor 2.2 package DEDS. For more details about the spike-in experiment, see http://www.affymetrix.com.

#### Usage

affySpikeIn

#### Format

affySpikeIn a numeric matrix of gene expression levels with 12626 rows (genes) and 24 columns (samples).

affySpikeIn.L a vector indicating the sample groups.

affySpikeIn.gnames a character vector containing the names of the genes.

spikedgene a numeric vector indicating the locations of the 14 spiked genes.

plot.ROTS

Plotting of a ROTS object

#### Description

Plots the ROTS objects created with the ROTS package.

### Usage

```
## S3 method for class 'ROTS'
plot(x, fdr=0.05, type="volcano", ...)
```

#### Arguments

Х	A ROTS object created from differential expression testing run by ROTS.
fdr	Selected cutoff for FDR value.
type	Type of plot to be created. Options are volcano (default) and heatmap.
	Other arguments passed to the plot function.

#### ROTS

#### Details

This function plots the results from a ROTS object using given false discovery rate threshold.

#### Value

Plots the results from ROTS object.

### Author(s)

Fatemeh Seyednasrollah, Tomi Suomi, Laura L. Elo Maintainer: Fatemeh Seyednasrollah <fatsey@utu.fi>

#### See Also

affySpikeIn

#### Examples

```
## ROTS-statistic for the Affymetrix spike-in data.
rots.out <- ROTS(data = affySpikeIn, groups = c(rep(0,5), rep(1,5)),
        B = 100, K = 500, seed = 1234)
## Plotting of the ROTS results.
plot(rots.out)</pre>
```

ROTS

*Reproducibility-Optimized Test Statistic (ROTS)* 

#### Description

Calculates the reproducibility-optimized test statistic (ROTS) for ranking genes in order of evidence for differential expression in two-group comparisons.

#### Usage

```
ROTS(data, groups, B, K, paired = FALSE, seed = NULL, a1 = NULL, a2 = NULL, log = TRUE, progress = FALSE
```

#### Arguments

data	a numeric data matrix or an ExpressionSet instance, in which rows correspond to genes and columns correspond to samples.
groups	a vector indicating the sample groups.
В	an integer specifying the number of bootstrap and permutation resamplings.
К	an integer indicating the largest top list size considered.
paired	a logical indicating whether a paired test is performed.
seed	an integer seed for the random number generator.
a1, a2	a non-negative parameter. See details section for further information.
log	a logical indicating whether input data is log2 scaled.
progress	a logical indicating if additional progress bars are shown.

#### Details

The reproducibility-optimization procedure ROTS enables the selection of a suitable gene ranking statistic directly from the given dataset. The statistic is optimized among a family of t-type statistics d = m/(a1+a2\*s), where m is the absolute difference between the group averages, s is the pooled standard error, and a1 and a2 are the non-negative parameters to be optimized. Two special cases of this family are the ordinary t-statistic (a1=0, a2=1) and the signal log-ratio (a1=1, a2=0). The optimality is defined in terms of maximal overlap of top-ranked genes in group-preserving bootstrap datasets. Importantly, besides the group labels, no a priori information about the properties of the data is required and no fixed cutoff for the gene rankings needs to be specified. For more details about the reproducibility-optimization procedure, see Elo et al. (2008).

The user is given the option to adjust the largest top list size considered in the reproducibility calculations, since lowering this size can markedly reduce the computation time. In large data matrices with thousands of rows, we generally recommend using a size of several thousands. In smaller data matrices, and especially if there are many rows with only a few non-missing entries, the size of K should be decreased accordingly.

ROTS tolerates a moderate number of missing values in the data matrix by effectively ignoring their contribution during the operation of the procedure. However, each row of the data matrix must contain at least two values in both groups. The rows containing only a few non-missing values should be removed; or alternatively, the missing data entries can be imputed using, e.g., the K-nearest neighbour imputation, which is implemented in the Bioconductor package impute.

If the parameter values a1 and a2 are set by the user, then no optimization is performed but the statistic and FDR-values are calculated for the given parameters. The false discovery rate (FDR) for the optimized test statistic is calculated by permuting the sample labels. The results for all the genes can be obtained by setting the FDR cutoff to 1.

#### Value

ROTS returns an object of class ROTS, which is a list containing the following components

data	the expression data matrix.
В	the number of bootstrap and permutation resamplings.
d	the value of the optimized ROTS-statistic for each gene.
pvalue	the corresponding pvalues.
FDR	the corresponding FDR-values.
a1	the optimized parameter a1.
a2	the optimized parameter a2.
k	the optimized top list size.
R	the optimized reproducibility value.
Z	the optimized reproducibility Z-score.

print prints the optimized parameters a1 and a2, the optimized top list size and the corresponding reproducibility values.

summary summarizes the results of a ROTS analysis. If fdr and num.genes are not specified, then the optimized parameters a1 and a2, the optimized top list size and the corresponding reproducibility values are shown. If fdr or num.genes is specified, then also the gene-specific information is shown for the genes at the specified FDR-level or top list size, respectively.

#### Author(s)

Fatemeh Seyednasrollah, Tomi Suomi, Laura L. Elo Maintainer: Fatemeh Seyednasrollah <fatsey@utu.fi>

#### References

L. L. Elo, S. Filen, R. Lahesmaa and T. Aittokallio: Reproducibility-optimized test statistic for ranking genes in microarray studies. IEEE/ACM Transactions on Computational Biology and Bioinformatics 5: 423–431, 2008.

#### See Also

affySpikeIn

#### Examples

```
## ROTS-statistic for the Affymetrix spike-in data.
rots.out <- ROTS(data = affySpikeIn, groups = c(rep(0,5), rep(1,5)),
        B = 100, K = 500, seed = 1234)
## Summary of the ROTS results.
rots.summary <- summary(rots.out, fdr = 0.05)</pre>
```

summary.ROTS Summary of a ROTS object

#### Description

Summarizes the differential testing results from ROTS package.

#### Usage

```
## S3 method for class 'ROTS'
summary(object, fdr=NULL, num.genes=NULL, verbose=TRUE, ...)
```

#### Arguments

object	a ROTS object created from differential expression testing run by ROTS.
fdr	selected cutoff for FDR value.
num.genes	selected cutoff number for number of differentially detected features.
verbose	If TRUE (default), summary function will print out 10 first detections which fulfill the cutoff criteria.
	other arguments passed to the summary function.

#### Details

This function returns the summary information (including row number, test-statistic, pvalue and FDR value) for the selected features.

#### Value

Returns a matrix where the rows are the selected features and columns are the Row number, ROTSstatistic, pvalue and FDR.

#### Author(s)

Fatemeh Seyednasrollah, Tomi Suomi, Laura L. Elo

Maintainer: Fatemeh Seyednasrollah <fatsey@utu.fi>

### References

L. L. Elo, S. Filen, R. Lahesmaa and T. Aittokallio: Reproducibility-optimized test statistic for ranking genes in microarray studies. IEEE/ACM Transactions on Computational Biology and Bioinformatics 5: 423–431, 2008.

#### See Also

affySpikeIn

#### Examples

```
## ROTS-statistic for the Affymetrix spike-in data.
rots.out <- ROTS(data = affySpikeIn, groups = c(rep(0,5), rep(1,5)),
        B = 100, K = 500 , seed = 1234)
## Summary of the ROTS results.
rots.summary <- summary(rots.out, fdr = 0.05)</pre>
```

upsSpikeIn

Protein expression data from the CPTAC Technology Assessment (Study 6)

#### Description

The data contains preprocessed expression values from CPTAC Technology Assessment, where equimolar concentrations of 48 human proteins (Sigma UPS 1) are spiked in five different concentrations into yeast proteome background and then processed. The example data contains only two groups from the study 6.

#### Usage

upsSpikeIn

#### Format

upsSpikeIn A numeric matrix of protein expressions.

## Index

\*Topic **datasets** affySpikeIn, 2 upsSpikeIn, 6 \*Topic **math** plot.ROTS, 2 ROTS, 3 summary.ROTS, 5

affySpikeIn, 2, 3, 5, 6

plot.ROTS, 2
print.ROTS(ROTS), 3

ROTS, 3 ROTS-package (ROTS), 3

spikedgene (affySpikeIn), 2
summary.ROTS, 5

upsSpikeIn,6