Package 'BLMA'

October 14, 2021

Date 2020-06-22
Type Package
Title BLMA: A package for bi-level meta-analysis
Version 1.16.0
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Description Suit of tools for bi-level meta-analysis. The package can be used in a wide range of applications, including general hypothesis testings, differential expression analysis, functional analysis, and pathway analysis.
biocViews GeneSetEnrichment, Pathways, DifferentialExpression, Microarray
License GPL (>=2)
Depends ROntoTools, GSA, PADOG, limma, graph, stats, utils, parallel, Biobase, metafor, methods
Suggests RUnit, BiocGenerics
RoxygenNote 7.1.0
NeedsCompilation no
git_url https://git.bioconductor.org/packages/BLMA
git_branch RELEASE_3_13
git_last_commit 9b4e0e3
git_last_commit_date 2021-05-19
Date/Publication 2021-10-14

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```
addCLT
```

The additive method for meta-analysis

Description

Combine independent studies using the average of p-values

Usage

addCLT(x)

Arguments

х

is an array of independent p-values

Details

This method is based on the fact that sum of independent uniform variables follow the Irwin-Hall distribution [1a,1b]. When the number of p-values is small (n < 20), the distribution of the average of p-values can be calculated using a linear transformation of the Irwin-Hall distribution. When n is large, the distribution is approximated using the Central Limit Theorem to avoid underflow/overflow problems [2,3,4,5].

Value

combined p-value

Author(s)

Tin Nguyen and Sorin Draghici

References

[1a] P. Hall. The distribution of means for samples of size n drawn from a population in which the variate takes values between 0 and 1, all such values being equally probable. Biometrika, 19(3-4):240-244, 1927.

[1b] J. O. Irwin. On the frequency distribution of the means of samples from a population having any law of frequency with finite moments, with special reference to Pearson's Type II. Biometrika, 19(3-4):225-239, 1927.

[2] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

[3] T. Nguyen, C. Mitrea, R. Tagett, and S. Draghici. DANUBE: Data-driven meta-ANalysis using UnBiased Empirical distributions – applied to biological pathway analysis. Proceedings of the IEEE, PP(99):1-20, 2016.

[4] T. Nguyen, D. Diaz, R. Tagett, and S. Draghici. Overcoming the matched-sample bottleneck: an orthogonal approach to integrate omic data. Scientific Reports, 6:29251, 2016.

[5] T. Nguyen, D. Diaz, and S. Draghici. TOMAS: A novel TOpology-aware Meta-Analysis approach applied to System biology. In Proceedings of the 7th ACM International Conference on Bioin- formatics, Computational Biology, and Health Informatics, pages 13-22. ACM, 2016.

See Also

fisherMethod, stoufferMethod

Examples

```
x <- rep(0,10)
addCLT(x)</pre>
```

x <- runif(10) addCLT(x)

bilevelAnalysisClassic

Bi-level meta-analysis in conjunction with a classical hypothesis testing method

Description

Perform a bi-level meta-analysis in conjunction with any of the classical hypothesis testing methods, such as t-test, Wilcoxon test, etc.

Usage

```
bilevelAnalysisClassic(x, y = NULL, splitSize = 5, metaMethod = addCLT,
func = t.test, p.value = "p.value", ...)
```

Arguments

х	a list of numeric vectors
У	an optional list of numeric vectors
splitSize	the minimum number of size in each split sample. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (additive method [1]), fishersMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])
func	the name of the hypothesis test. By default func=t.test
p.value	the component that returns the p-value after performing the test provided by the <i>func</i> parameter. For example, the function t-test returns the class "htest" where the component "p.value" is the p-value of the test. By default, p.value="p.value"
	additional parameters for <i>func</i>

Details

This function performs a bi-level meta-analysis for the lists of samples [1]. It performs intraexperiment analyses to compare the vectors in x agains the corresponding vectors in y using the function intraAnalysisClassic in conjunction with the test provided in *func*. For example, it compares the first vector in x with the first vector in y, the second vector in x with the second vector in y, etc. When y is null, then the comparisons are reduced to one-sample tests. After these comparisons, we have a list of p-values, one for each comparision. The function then combines these p-values to obtain a single p-value using *metaMethod*.

Value

the combined p-value

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

See Also

intraAnalysisClassic, intraAnalysisGene, bilevelAnalysisGene

Examples

```
set.seed(1)
11 <- lapply(as.list(seq(3)),FUN=function (x) rnorm(n=10, mean=1))
11
# one-sample t-test
lapply(11, FUN=function(x) t.test(x, alternative="greater")$p.value)</pre>
```

```
# combining the p-values of one-sample t-tests:
addCLT(unlist(lapply(l1, FUN=function(x) t.test(x, alter="g")$p.value)))
#Bi-level meta-analysis
bilevelAnalysisClassic(x=l1, alternative="greater")
```

bilevelAnalysisGene Bi-level meta-analysis of multiple expression datasets at the gene-level

Description

Perform a bi-level meta-analysis in conjunction with the moderate t-test (limma package) for the purpose of differential expression analysis of multiple gene expression datasets

Usage

```
bilevelAnalysisGene(dataList, groupList, splitSize = 5, metaMethod = addCLT)
```

Arguments

dataList	a list of datasets. Each dataset is a data frame where the rows are the gene IDs and the columns are the samples
groupList	a list of vectors. Each vector represents the phenotypes of the corresponding dataset in dataList, which are either 'c' (control) or 'd' (disease).
splitSize	the minimum number of disease samples in each split dataset. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (additive method [1]), fishersMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])

Details

The bi-level framework combines the datasets at two levels: an intra- experiment analysis, and an inter-experiment analysis [1]. At the intra-experiment analysis, the framework splits a dataset into smaller datasets, performs a moderated t-test (limma package) on split datasets, and then combines p-values of individual genes using *metaMethod*. At the inter-experiment analysis, the p-values obtained for each individual datasets are combined using *metaMethod*

Value

A data frame containing the following components:

- rownames: gene IDs that are common in all datasets
- pLimma: the p-values obtained by combining pLimma values of individual datasets
- pLimma.fdr: FDR-corrected p-values of pLimma
- pBilevel: the p-values obtained from combining pIntra values of individual datasets
- *pBilevel.fdr:* FDR-corrected p-values of pBilevel

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

See Also

bilevelAnalysisGene, intraAnalysisClassic

Examples

```
dataSets <- c("GSE17054", "GSE57194", "GSE33223", "GSE42140")
data(list=dataSets, package="BLMA")
names(dataSets) <- dataSets
dataList <- lapply(dataSets, function(dataset) get(paste0("data_", dataset)))
groupList <- lapply(dataSets, function(dataset) get(paste0("group_", dataset)))
Z <- bilevelAnalysisGene(dataList = dataList, groupList = groupList)
head(Z)</pre>
```

bilevelAnalysisGeneset

Bi-level meta-analysis – applied to geneset enrichment analysis

Description

Perform a bi-level meta-analysis in conjunction with geneset enrichment methods (ORA/GSA/PADOG) to integrate multiple gene expression datasets.

Usage

```
bilevelAnalysisGeneset(gslist, gs.names, dataList, groupList, splitSize = 5,
metaMethod = addCLT, enrichment = "ORA", pCutoff = 0.05,
percent = 0.05, mc.cores = 1, ...)
```

Arguments

gslist	a list of gene sets.
gs.names	names of the gene sets.
dataList	a list of datasets to be combined. Each dataset is a data frame where the rows are the gene IDs and the columns are the samples.
groupList	a list of vectors. Each vector represents the phenotypes of the corresponding dataset in dataList. The elements of each vector are either 'c' (control) or 'd' (disease).

splitSize	the minimum number of disease samples in each split dataset. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (addi- tive method [1]), fisherMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])
enrichment	the method used for enrichment analysis. This should be one of "ORA", "GSA", or "PADOG". By default, enrichment is set to "ORA".
pCutoff	cutoff p-value used to identify differentially expressed (DE) genes. This parameter is used only when the enrichment method is "ORA". By default, pCut-off=0.05 (five percent)
percent	percentage of genes with highest foldchange to be considered as differentially expressed (DE). This parameter is used when the enrichment method is "ORA". By default percent=0.05 (five percent). Please note that only genes with p-value less than pCutoff will be considered
mc.cores	the number of cores to be used in parallel computing. By default, mc.cores=1
	additional parameters of the GSA/PADOG functions

Details

The bi-level framework combines the datasets at two levels: an intra- experiment analysis, and an inter-experiment analysis [1]. At the intra-level analysis, the framework splits a dataset into smaller datasets, performs enrichment analysis for each split dataset (using ORA [2], GSA [3], or PADOG [4]), and then combines the results of these split datasets using *metaMethod*. At the inter-level analysis, the results obtained for individual datasets are combined using *metaMethod*

Value

A data frame (rownames are geneset/pathway IDs) that consists of the following information:

- Name: name/description of the corresponding pathway/geneset
- Columns that include the pvalues obtained from the intra-experiment analysis of individual datasets
- pBLMA: p-value obtained from the inter-experiment analysis using addCLT
- rBLMA: ranking of the geneset/pathway using addCLT
- pBLMA.fdr: FDR-corrected p-values

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

[2] S. Draghici, P. Khatri, R. P. Martin, G. C. Ostermeier, and S. A. Krawetz. Global functional profiling of gene expression. Genomics, 81(2):98-104, 2003.

[3] B. Efron and R. Tibshirani. On testing the significance of sets of genes. The Annals of Applied Statistics, 1(1):107-129, 2007.

[4] A. L. Tarca, S. Draghici, G. Bhatti, and R. Romero. Down-weighting overlapping genes improves gene set analysis. BMC Bioinformatics, 13(1):136, 2012.

[5] R. A. Fisher. Statistical methods for research workers. Oliver & Boyd, Edinburgh, 1925.

[6] S. Stouffer, E. Suchman, L. DeVinney, S. Star, and J. Williams, RM. The American Soldier: Adjustment during army life, volume 1. Princeton University Press, Princeton, 1949.

[7] L. H. C. Tippett. The methods of statistics. The Methods of Statistics, 1931.

[8] B. Wilkinson. A statistical consideration in psychological research. Psychological Bulletin, 48(2):156, 1951.

See Also

bilevelAnalysisPathway, phyper, GSA, padog

Examples

```
# load KEGG pathways and create gene sets
x <- loadKEGGPathways()</pre>
gslist <- lapply(x$kpg,FUN=function(y){return (nodes(y));})</pre>
gs.names <- x$kpn[names(gslist)]</pre>
# load example data
dataSets <- c("GSE17054", "GSE57194", "GSE33223", "GSE42140")</pre>
data(list=dataSets, package="BLMA")
names(dataSets) <- dataSets</pre>
dataList <- lapply(dataSets, function(dataset) get(paste0("data_", dataset)))</pre>
groupList <- lapply(dataSets, function(dataset) get(paste0("group_", dataset)))</pre>
# perform bi-level meta-analysis in conjunction with ORA
ORAComb <- bilevelAnalysisGeneset(gslist, gs.names, dataList, groupList, enrichment = "ORA")
head(ORAComb[, c("Name", "pBLMA", "pBLMA.fdr", "rBLMA")])
# perform bi-level meta-analysis in conjunction with GSA
GSAComb <- bilevelAnalysisGeneset(gslist, gs.names, dataList, groupList, enrichment = "GSA", nperms = 200, random.
head(GSAComb[, c("Name", "pBLMA", "pBLMA.fdr", "rBLMA")])
# perform bi-level meta-analysi in conjunction with PADOG
set.seed(1)
```

```
PADOGComb <- bilevelAnalysisGeneset(gslist, gs.names, dataList, groupList, enrichment = "PADOG", NI=200)
head(PADOGComb[, c("Name", "pBLMA", "pBLMA.fdr", "rBLMA")])
```

bilevelAnalysisPathway

Bi-level meta-analysis – applied to pathway analysis

Description

Perform a bi-level meta-analysis conjunction with Impact Analysis to integrate multiple gene expression datasets

Usage

```
bilevelAnalysisPathway(kpg, kpn, dataList, groupList, splitSize = 5,
metaMethod = addCLT, pCutoff = 0.05, percent = 0.05, mc.cores = 1,
nboot = 200, seed = 1)
```

Arguments

kpg	list of pathway graphs as objects of type graph (e.g., graphNEL)
kpn	names of the pathways.
dataList	a list of datasets to be combined. Each dataset is a data frame where the rows are the gene IDs and the columns are the samples.
groupList	a list of vectors. Each vector represents the phenotypes of the corresponding dataset in dataList, which are either 'c' (control) or 'd' (disease).
splitSize	the minimum number of disease samples in each split dataset. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (addi- tive method [1]), fisherMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])
pCutoff	cutoff p-value used to identify differentially expressed (DE) genes. This parameter is used only when the enrichment method is "ORA". By default, pCut-off=0.05 (five percent)
percent	percentage of genes with highest foldchange to be considered as differentially expressed (DE). This parameter is used when the enrichment method is "ORA". By default percent=0.05 (five percent). Please note that only genes with p-value less than pCutoff will be considered
mc.cores	the number of cores to be used in parallel computing. By default, mc.cores=1
nboot	number of bootstrap iterations. By default, nboot=200
seed	seed. By default, seed=1.

Details

The bi-level framework combines the datasets at two levels: an intra-experiment analysis, and an inter-experiment analysis [1]. At the intra-level analysis, the framework splits a dataset into smaller datasets, performs pathway analysis for each split dataset using Impact Analysis [2,3], and then combines the results of these split datasets using *metaMethod*. At the inter-level analysis, the results obtained for individual datasets are combined using *metaMethod*

Value

A data frame (rownames are geneset/pathway IDs) that consists of the following information:

- Name: name/description of the corresponding pathway/geneset
- Columns that include the pvalues obtained from the intra-experiment analysis of individual datasets
- pBLMA: p-value obtained from the inter-experiment analysis using addCLT
- rBLMA: ranking of the geneset/pathway using addCLT
- *pBLMA.fdr:* FDR-corrected p-values

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

[2] A. L. Tarca, S. Draghici, P. Khatri, S. S. Hassan, P. Mittal, J.-s. Kim, C. J. Kim, J. P. Kusanovic, and R. Romero. A novel signaling pathway impact analysis. Bioinformatics, 25(1):75-82, 2009.

[3] S. Draghici, P. Khatri, A. L. Tarca, K. Amin, A. Done, C. Voichita, C. Georgescu, and R. Romero. A systems biology approach for pathway level analysis. Genome Research, 17(10):1537-1545, 2007.

[4] R. A. Fisher. Statistical methods for research workers. Oliver & Boyd, Edinburgh, 1925.

[5] S. Stouffer, E. Suchman, L. DeVinney, S. Star, and J. Williams, RM. The American Soldier: Adjustment during army life, volume 1. Princeton University Press, Princeton, 1949.

[6] L. H. C. Tippett. The methods of statistics. The Methods of Statistics, 1931.

[7] B. Wilkinson. A statistical consideration in psychological research. Psychological Bulletin, 48(2):156, 1951.

See Also

bilevelAnalysisGeneset, pe, phyper

Examples

```
# load KEGG pathways
x <- loadKEGGPathways()
# load example data
dataSets <- c("GSE17054", "GSE57194", "GSE33223", "GSE42140")
data(list=dataSets, package="BLMA")
names(dataSets) <- dataSets
dataList <- lapply(dataSets, function(dataset) get(paste0("data_", dataset)))
groupList <- lapply(dataSets, function(dataset) get(paste0("group_", dataset)))</pre>
```

IAComb <- bilevelAnalysisPathway(x\$kpg, x\$kpn, dataList, groupList)</pre>

fisherMethod

```
head(IAComb[, c("Name", "pBLMA", "pBLMA.fdr", "rBLMA")])
```

fisherMethod Fisher's method for meta-analysis

Description

Combine independent p-values using the minus log product

Usage

fisherMethod(x)

Arguments ×

is an array of independent p-values

Details

Considering a set of m independent significance tests, the resulted p-values are independent and uniformly distributed between 0 and 1 under the null hypothesis. Fisher's method uses the minus log product of the p-values as the summary statistic, which follows a chi-square distribution with 2m degrees of freedom. This chi-square distribution is used to calculate the combined p-value.

Value

combined p-value

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] R. A. Fisher. Statistical methods for research workers. Oliver & Boyd, Edinburgh, 1925.

See Also

stoufferMethod, addCLT

Examples

```
x <- rep(0,10)
fisherMethod(x)</pre>
```

x <- runif(10)
fisherMethod(x)</pre>

getStatistics

Description

Calculate genes summary statistic across multiple datasets

Usage

```
getStatistics(allGenes, dataList, groupList, ncores = 1, method = addCLT)
```

Arguments

allGenes	Vector of all genes names for the analysis.
dataList	A list of expression matrices, in which rows are genes and columns are samples.
groupList	A list of vectors indicating sample group corresponding with expression matrices in dataList.
ncores	Number of core to use in prallel processing.
method	Function for combining p-values. It must accept one input which is a vector of p-values and return a combined p-value. Three methods are embedded in this package are addCLT, fisherMethod, and stoufferMethod.

Details

To estimate the effect sizes of genes across all studies, first standardized mean difference for each gene in individual studies is compute. Next, the overall effect size and standard error are estimated using the random-effects model. This overall effect size represents the gene's expression change under the effect of the condition. The, z-scores and p-values of observing such effect sizes are computed. The p-values is obtained from classical hypothesis testing. By default, linear model and empirical Bayesian testing \(limma\) are used to compute the p-values for differential expression. The two-tailed p-values are converted to one-tailed p-values (lef- and right-tailed). For each gene, the one-tailed p-values across all datasets are then combined using the addCLT, stouffer or fisher method. These p-values represent how likely the differential expression is observed by chance.

Value

A data.frame of gene statistics with following columns:

pTwoTails Two-tailed p-values **pTwoTails.fdr** Two-tailed p-values with false discovery rate correction **pLeft** left-tailed p-values **pLeft.fdr** left-tailed p-values with false discovery rate correction **pRight.fdr** right-tailed p-values with false discovery rate correction **pRight** right-tailed p-values

getStatistics

ES Effect size

ES.pTwoTails Two-tailed p-values for effect size

ES.pTwoTails.fdr Two-tailed p-values for effect size with false discovery rate correction

ES.pLeft Left-tailed p-values for effect size

ES.pLeft.fdr Left-tailed p-values for effect size with false discovery rate correction

ES.pRight Right-tailed p-values for effect size

ES.pRight.fdr Right-tailed p-values for effect size with false discovery rate correction

Author(s)

Tin Nguyen, Hung Nguyen, and Sorin Draghici

References

Nguyen, T., Shafi, A., Nguyen, T. M., Schissler, A. G., & Draghici, S. (2020). NBIA: a networkbased integrative analysis framework-applied to pathway analysis. Scientific reports, 10(1), 1-11. Nguyen, T., Tagett, R., Donato, M., Mitrea, C., & Draghici, S. (2016). A novel bi-level metaanalysis approach: applied to biological pathway analysis. Bioinformatics, 32(3), 409-416. Smyth, G. K. (2005). Limma: linear models for microarray data. In Bioinformatics and computational biology solutions using R and Bioconductor (pp. 397-420). Springer, New York, NY.

See Also

addCLT

Examples

```
datasets <- c("GSE17054", "GSE57194", "GSE33223", "GSE42140")</pre>
data(list = datasets, package = "BLMA")
dataList <- lapply(datasets, function(dataset) {</pre>
    get(paste0("data_", dataset))
})
groupList <- lapply(datasets, function(dataset) {</pre>
    get(paste0("group_", dataset))
})
names(dataList) <- datasets</pre>
names(groupList) <- datasets</pre>
allGenes <- Reduce(intersect, lapply(dataList, rownames))
geneStat <- getStatistics(allGenes, dataList, groupList)</pre>
head(geneStat)
# perform pathway analysis
library(ROntoTools)
# get gene network
kpg <- loadKEGGPathways()$kpg</pre>
# get gene network name
kpn <- loadKEGGPathways()$kpn</pre>
```

```
# get geneset
gslist <- lapply(kpg,function(y) nodes(y))</pre>
# get differential expressed genes
DEGenes.Left <- rownames(geneStat)[geneStat$pLeft < 0.05 & geneStat$ES.pLeft < 0.05]</pre>
DEGenes.Right <- rownames(geneStat)[geneStat$pRight < 0.05 & geneStat$ES.pRight < 0.05]</pre>
DEGenes <- union(DEGenes.Left, DEGenes.Right)</pre>
# perform pathway analysis with ORA
oraRes <- lapply(gslist, function(gs){</pre>
    pORACalc(geneSet = gs, DEGenes = DEGenes, measuredGenes = rownames(geneStat))
})
oraRes <- data.frame(p.value = unlist(oraRes), pathway = names(oraRes))</pre>
rownames(oraRes) <- kpn[rownames(oraRes)]</pre>
# print results
print(head(oraRes))
# perfrom pathway analysis with Pathway-Express from ROntoTools
ES <- geneStat[DEGenes, "ES"]</pre>
names(ES) <- DEGenes</pre>
peRes = pe(x = ES, graphs = kpg, ref = allGenes, nboot = 1000, seed=1)
peRes.Summary <- Summary(peRes, comb.pv.func = fisherMethod)</pre>
peRes.Summary[, ncol(peRes.Summary) + 1] <- rownames(peRes.Summary)</pre>
rownames(peRes.Summary) <- kpn[rownames(peRes.Summary)]</pre>
colnames(peRes.Summary)[ncol(peRes.Summary)] = "pathway"
# print results
print(head(peRes.Summary))
```

GSE17054

Gene expression dataset GSE17054 from Majeti et al.

Description

This dataset consists of 5 acute myeloid leukemia and 4 control samples. The data frame data_GSE17054 includes the expression data while the vector group_GSE17054 includes the grouping information.

Usage

```
data(GSE17054)
```

Format

data_GSE17054 is a data frame with 4738 rows and 9 columns. The rows represent the genes and the columns represent the samples.

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GSE33223

group_GSE17054 is a vector that represents the sample grouping for data_GSE17054. The elements of *group_GSE17054* are either 'c' (control) or 'd' (disease).

Source

Obtained from http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE17054

References

Majeti et al. Dysregulated gene expression networks in human acute myelogenous leukemia stem cells. Proceedings of the National Academy of Sciences, 106(9):3396-3401, 2009.

GSE33223

Gene expression dataset GSE33223 from Bacher et al.

Description

This dataset consists of 20 acute myeloid leukemia and 10 control samples. The data frame data_GSE33223 includes the expression data while the vector group_GSE33223 includes the grouping information.

Usage

data(GSE33223)

Format

data_GSE33223 is a data frame with 4114 rows and 30 columns. The rows represent the genes and the columns represent the samples.

group_GSE33223 is a vector that represents the sample grouping for data_GSE33223. The elements of *group_GSE33223* are either 'c' (control) or 'd' (disease).

Source

Obtained from https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE33223

References

Bacher et al. Multilineage dysplasia does not influence prognosis in CEBPA-mutated AML, supporting the WHO proposal to classify these patients as a unique entity. Blood, 119(20):4719-22, 2012.

GSE42140

Description

This dataset consists of 26 acute myeloid leukemia and 5 control samples. The data frame data_GSE42140 includes the expression data while the vector group_GSE42140 includes the grouping information.

Usage

data(GSE42140)

Format

data_GSE42140 is a data frame with 4114 rows and 31 columns. The rows represent the genes and the columns represent the samples.

group_GSE42140 is a vector that represents the sample grouping for data_GSE42140. The elements of *group_GSE42140* are either 'c' (control) or 'd' (disease).

References

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE42140

GSE57194 Gene expression dataset GSE57194 from Abdul-Nabi et al.	
--	--

Description

This dataset consists of 6 acute myeloid leukemia and 6 control samples. The data frame data_GSE57194 includes the expression data while the vector group_GSE57194 includes the grouping information.

Usage

data(GSE57194)

Format

data_GSE57194 is a data frame with 4114 rows and 12 columns. The rows represent the genes and the columns represent the samples.

group_GSE57194 is a vector that represents the sample grouping for data_GSE57194. The elements of *group_GSE57194* are either 'c' (control) or 'd' (disease).

Source

Obtained from https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE57194

intraAnalysisClassic

References

Abdul-Nabi et al. In vitro transformation of primary human CD34+ cells by AML fusion oncogenes: early gene expression profiling reveals possible drug target in AML. PLoS One, 5(8):e12464, 2010.

intraAnalysisClassic Intra-experiment analysis in conjunction with classical hypothesis tests

Description

Perform an intra-experiment analysis in conjunction with any of the classical hypothesis testing methods, such as t-test, Wilcoxon test, etc.

Usage

```
intraAnalysisClassic(x, y = NULL, splitSize = 5, metaMethod = addCLT,
func = t.test, p.value = "p.value", ...)
```

Arguments

х	a numeric vector of data values
У	an optional numeric vector of values
splitSize	the minimum number of size in each split sample. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (additive method [1]), fishersMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])
func	the name of the hypothesis test. By default func=t.test
p.value	the component that returns the p-value after performing the test provided by the <i>func</i> parameter. For example, the function t-test returns the class "htest" where the component "p.value" is the p-value of the test. By default, p.value="p.value"
	additional parameters for <i>func</i>

Details

This function performs an intra-experiment analysis for the given sample(s) [1]. Given x as the numeric vector, this function first splits x into smaller samples with size *splitSize*, performs hypothesis testing using *func*, and then combines the p-values using *metaMethod*

Value

intra-experiment p-value

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

See Also

bilevelAnalysisClassic, intraAnalysisGene, bilevelAnalysisGene

Examples

```
set.seed(1)
x <- rnorm(10, mean = 0)
# p-value obtained from a one-sample t-test
t.test(x, mu=1, alternative = "less")$p.value
# p-value obtained from an intra-experiment analysis
intraAnalysisClassic(x, func=t.test, mu=1, alternative = "less")
# p-value obtained from a one-sample wilcoxon test
wilcox.test(x, mu=1, alternative = "less")$p.value
# p-value obtained from an intra-experiment analysis
intraAnalysisClassic(x, func=wilcox.test, mu=1, alternative = "less")
set.seed(1)
x <- rnorm(20, mean=0); y <- rnorm(20, mean=1)</pre>
# p-value obtained from a two-sample t-test
t.test(x,y,alternative="less")$p.value
# p-value obtained from an intra-experiment analysis
intraAnalysisClassic(x, y, func=t.test, alternative = "less")
# p-value obtained from a two-sample wilcoxon test
wilcox.test(x,y,alternative="less")$p.value
# p-value obtained from an intra-experiment analysis
intraAnalysisClassic(x, y, func=wilcox.test, alternative = "less")
```

intraAnalysisGene Intra-experiment analysis of an expression dataset at the gene-level

Description

perform an intra-experiment analysis in conjunction with the moderated t-test (limma package) for the purpose of differential expression analysis of a gene expression dataset

Usage

```
intraAnalysisGene(data, group, splitSize = 5, metaMethod = addCLT)
```

intraAnalysisGene

Arguments

data	a data frame where the rows are the gene IDs and the columns are the samples
group	sample grouping. The elements of <i>group</i> are either 'c' (control) or 'd' (disease). names(group) should be identical to colnames(data)
splitSize	the minimum number of disease samples in each split dataset. splitSize should be at least 3. By default, splitSize=5
metaMethod	the method used to combine p-values. This should be one of addCLT (additive method [1]), fishersMethod (Fisher's method [5]), stoufferMethod (Stouffer's method [6]), max (maxP method [7]), or min (minP method [8])

Details

This function performs an intra-experiment analysis [1] for individual genes of the given dataset. The function first splits the dataset into smaller datasets, performs a moderated t-test (limma pack-age) for the genes of the split datasets, and then combines the p-values for individual genes using *metaMethod*

Value

A data frame (rownames are gene IDs) that consists of the following information:

- *logFC*: log foldchange (diseases versus controls)
- pLimma: p-value obtained from limma without spliting
- pLimma.fdr: FDR-corrected p-values of pLimma
- pIntra: p-value obtained from intra-experiment analysis
- pIntra.fdr: FDR-corrected p-values of pIntra

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] T. Nguyen, R. Tagett, M. Donato, C. Mitrea, and S. Draghici. A novel bi-level meta-analysis approach – applied to biological pathway analysis. Bioinformatics, 32(3):409-416, 2016.

See Also

bilevelAnalysisGene, intraAnalysisClassic, link{bilevelAnalysisClassic}

Examples

```
data(GSE33223)
X <- intraAnalysisGene(data_GSE33223, group_GSE33223)
head(X)</pre>
```

loadKEGGPathways Load KEGG pathways and names

Description

Load KEGG pathways and names

Usage

```
loadKEGGPathways(organism = "hsa", updateCache = FALSE)
```

Arguments

organism	organism code. Default value is "hsa" (human)	
updateCache	re-download KEGG pathways. Default value is FALSE	

Value

A list of the following components

- *kpg* a list of graphNEL objects encoding the pathway information.
- *kpn* a named vector of pathway tiles. The names of the vector are the pathway KEGG IDs.

Author(s)

Tin Nguyen and Sorin Draghici

See Also

keggPathwayGraphs, keggPathwayNames

Examples

x <- loadKEGGPathways()</pre>

pORACalc

Description

Calculate p-value for over-representation Analysis

Usage

pORACalc(geneSet, DEGenes, measuredGenes, minSize = 0)

Arguments

geneSet	a vector of gene names belong to the geneset	
DEGenes	a vector of differential expressed genes	
measuredGenes	a vector of all genes in the analysis	
minSize	the minimum number of DE genes in the geneSet	

Value

p-value

stoufferMethod	Stouffer's method for meta-analysis	

Description

Combine independent studies using the sum of p-values transformed into standard normal variables

Usage

stoufferMethod(x)

Arguments

х

is an array of independent p-values

Details

Considering a set of m independent significance tests, the resulted p-values are independent and uniformly distributed between 0 and 1 under the null hypothesis. Stouffer's method is similar to Fisher's method (fisherMethod), with the difference is that it uses the sum of p-values transformed into standard normal variables instead of the log product.

Value

combined p-value

Author(s)

Tin Nguyen and Sorin Draghici

References

[1] S. Stouffer, E. Suchman, L. DeVinney, S. Star, and R. M. Williams. The American Soldier: Adjustment during army life, volume 1. Princeton University Press, Princeton, 1949.

See Also

fisherMethod, addCLT

Examples

x <- rep(0,10)
stoufferMethod(x)</pre>

x <- runif(10)
stoufferMethod(x)</pre>

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