Package 'metapone'

October 15, 2023

Type Package

Title Conducts pathway test of metabolomics data using a weighted permutation test

Version 1.6.0

Date 2022-8-20

- **Description** The package conducts pathway testing from untargetted metabolomics data. It requires the user to supply feature-level test results, from case-control testing, regression, or other suitable feature-level tests for the study design. Weights are given to metabolic features based on how many metabolites they could potentially match to. The package can combine positive and negative mode results in pathway tests.
- **Depends** R (>= 4.1.0), BiocParallel, fields, markdown, fdrtool, fgsea, ggplot2, ggrepel

Imports methods

biocViews Technology, MassSpectrometry, Metabolomics, Pathways

License Artistic-2.0

LazyLoad yes

NeedsCompilation no

Suggests rmarkdown, knitr

VignetteBuilder knitr

git_url https://git.bioconductor.org/packages/metapone

git_branch RELEASE_3_17

git_last_commit bb8611a

git_last_commit_date 2023-04-25

Date/Publication 2023-10-15

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metapone-package

Conducts pathway test of metabolomics data using a weighted permutation test

Description

The package conducts pathway testing from untargetted metabolomics data. It requires the user to supply feature-level test results, from case-control testing, regression, or other suitable feature-level tests for the study design. Weights are given to metabolic features based on how many metabolites they could potentially match to. The package can combine positive and negative mode results in pathway tests. The package contains two types of statistical testing that considers matching uncertainty - (1) a permutation test that is based on the hypergeometric test and (2) a GSEA type test with weighted features/metabolites.

Details

The package conducts (1) a weighted hypergeometric test using permutations on metabolomics data. The weights are assigned based on how many metabolites each data feature can match to, (2) a GSEA type test based on an estimation of importance of metabolites/features. The importance is evluated by the size of matching for each metabolite/feature and the p-value of features.

The user can tune a parameter to change the penalty for multiple-matched features and choose the type of pathway testing.

Author(s)

Tianwei Yu (<yutianwei@cuhk.edu.cn>)

bbplot1d

Plot of metapone result.

Description

The function bbplot1d() select important pathways with their P-value less than a threshold and returns ranked bubble plot showing important pathways names and their corresponding -log10(Pvalue).

Usage

```
bbplot1d(res, p_thres = 0.05, sig_metab_thres = 1, low.color = "MidnightBlue", high.color = "LightSkyBl
```

Arguments

res	The result matrix obtained from metapone with columns: "p_value", "n_significant metabolites", "n_mapped_metabolites", "n_metabolites", "significant metabolites", "mapped_metabolites", "fdr".	
p_thres	The threshold of P-value for pathways to be shown in the bubble plot. The default threshold is 0.05.	
sig_metab_thres		
	The threshold of fractional matched significant metabolite count for pathways to be shown in the bubble plot. The default is 1.	
low.color	The GRB color of the lowest ldfr value to be shown in the bubble plot.	
high.color	The GRB color of the highest ldfr value to be shown in the bubble plot.	

Author(s)

LeqiTian(<leqitian@link.cuhk.edu.cn>)

See Also

metapone

```
data(hmdbCompMZ.metapone)
data(pa)
data(pos)
data(neg)
dat <- list(pos, neg)
type <- list("pos", "neg")
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone, p.threshold=0.05,
    n.permu=100,fractional.count.power=0.5, max.match.count=10)
bbplot1d(ptable(r)) # p_thres = 0.05</pre>
```

bbplot2d

Description

The function bbplot2d() select important pathways with their P-value less than a threshold and returns a 2-D bubble plot with -log10(Pvalue) and the number of significant metabolites as coordinate axes.

Usage

```
bbplot2d(res, p_thres = 0.05, sig_metab_thres = 1, low.color = "MidnightBlue", high.color = "LightSkyBl
```

Arguments

res	The result matrix obtained from metapone with columns: "p_value", "n_significant metabolites", "n_mapped_metabolites", "n_metabolites", "significant metabolites", "mapped_metabolites", "fdr".	
p_thres	The threshold of P-value for pathways to be shown in the bubble plot. The default threshold is 0.05.	
sig_metab_thres		
	The threshold of fractional matched significant metabolite count for pathways to be shown in the bubble plot. The default is 1.	
low.color	The GRB color of the lowest ldfr value to be shown in the bubble plot.	
high.color	The GRB color of the highest ldfr value to be shown in the bubble plot.	

Author(s)

LeqiTian(<leqitian@link.cuhk.edu.cn>)

See Also

metapone

```
data(hmdbCompMZ.metapone)
data(pa)
data(pos)
data(neg)
dat <- list(pos, neg)
type <- list("pos", "neg")
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone, p.threshold=0.05,
    n.permu=100,fractional.count.power=0.5, max.match.count=10)
bbplot2d(ptable(r)) # p_thres = 0.05</pre>
```

ftable

Description

Returns a list containing the mapped features in each pathway.

Usage

```
## S4 method for signature 'metaponeResult'
ftable(object)
```

Arguments

object A metaponeResult object.

Details

Each pathway is represented by a data.frame as an item in the list object. The dataframe include information of m.z, retention.time, p.value, statistic, HMDB_ID, theoretical m.z, ion.type, fractional counts.

Value

The method returns a list. Each item is for a pathway. Matched significant metabolites are included.

Author(s)

Tianwei Yu <yutianwei@cuhk.edu.cn>

See Also

ptable

```
data(hmdbCompMZ.metapone)
data(pa)
data(pos)
data(neg)
dat <- list(pos, neg)
type <- list("pos", "neg")
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone, p.threshold=0.05,
    n.permu=100,fractional.count.power=0.5, max.match.count=10)
ftable(r)[1:6]</pre>
```

hmdbCompMZ

Description

Monoisotopic mass of common adduct ions.

Usage

data("hmdbCompMZ")

Format

A data frame with 5704350 observations on the following 3 variables.

HMDB_ID HMDB ID.

ion.type Adduct ion type.

m.z the m/z of the adduct ion.

Source

https://hmdb.ca/

References

https://hmdb.ca/

Examples

data(hmdbCompMZ)

hmdbCompMZ.metapone the m/z values of common adduct ions of metapone metaboites

Description

Monoisotopic mass of common adduct ions, limited to those included in the pathways in metapone.

Usage

data("hmdbCompMZ.metapone")

metapone

Format

A data frame with 79350 observations on the following 3 variables.

HMDB_ID HMDB ID.

ion.type Adduct ion type.

m.z the m/z of the adduct ion.

Details

The main difference of using this dataset vs using hmdbCompMZ, is the metabolite universe in testing is limited to those metabolites matched to metapone pathways, not all HMDB metabolites.

Source

The Human Metabolome Database

References

The Human Metabolome Database

Examples

data(hmdbCompMZ)

metapone

METAbolic pathway testing using both POsitive and NEgative mode data

Description

Metapone conducts pathway tests for untargeted metabolomics data. It has three main characteristics: (1) expanded database combining SMPDB and Mummichog databases, with manual cleaning to remove redundancies; (2) A new weighted testing scheme to address the issue of metabolitefeature matching uncertainties; (3) Can consider positive mode and negative mode data in a single analysis.

Usage

```
metapone(dat=NULL, type=NULL, pa, hmdbCompMZ, pos.adductlist = c("M+H",
"M+NH4", "M+Na", "M+ACN+H","M+ACN+Na", "M+2ACN+H", "2M+H", "2M+Na",
"2M+ACN+H"), neg.adductlist = c("M-H","M-2H","M-2H+Na","M-2H+K",
"M-2H+NH4","M-H2O-H","M-H+Cl", "M+Cl", "M+2Cl"),
use.fractional.count=TRUE, match.tol.ppm=5, p.threshold=0.05,
n.permu=200, fractional.count.power=0.5, max.match.count=10,
use.fgsea = FALSE, use.meta = FALSE)
```

Arguments

	dat	The list of test results. An element in the list should be postive ion mode test re- sults or negative ion mode test results with four columns: m/z, retention time, p- value, test statistic. The package doesn't require both pos and neg to be present. One ion mode result is sufficient. Multiple ion mode results are allowed.
	type	The list of corresponding ion mode of each element in dat. Each element in the list should be "pos" or "neg". The size of type should be consistent with the size of dat.
	ра	Pathway information. A data frame with five columns: database pathway ID, pathway name, HMDB ID, KEGG ID, category of pathway.
	hmdbCompMZ	the m/z values of common adduct ions of HMDB metaboites. See the help file of hmdbCompMZ for details.
	<pre>pos.adductlist</pre>	The vector of positive adduct ions to be considered.
	neg.adductlist	The vector of negative adduct ions to be considered.
use.fractional.count		count
		A lot of features match to multiple metabolites by m/z. Whether to discount such matches by using fractional counts.
	<pre>match.tol.ppm</pre>	The ppm level when conducting m/z match.
	p.threshold	The threshold of p-values of metabolic features to be considered significant.
	n.permu	The number of permutations in permutation test.
	fractional.cour	nt.power
		The fractional counts are taken to this power to transform the weights.
	<pre>max.match.count</pre>	
		When calculating fractional counts, some features might be matched to too many. In that case the number of matches is capped by the value of max.match.count.
	use.fgsea	Whether to use a GSEA type test when performing pathway testing. When it is FALSE, a permutation-based weighted hypergeometric test is performed.
	use.meta	Whether to perform a GSEA type test with weighted metabolites. When it is FALSE, a GSEA type test is performed on weighted features.

Value

The method returns a generic S4 object of class "metapone.result":

@test.results	A matrix with 8 columns: "p_value", "n_significant metabolites", "n_mapped_metabolites",
	"n_metabolites", "significant metabolites", "mapped_metabolites", "lfdr", "ad-
	just.p". Each row is for a pathway. When using GSEA test, "ES", "NES",
	"nMoreExtreme" are returned additionally.

@mapped.features

A list. Each item is for a pathway. The item lists matched significant metabolites.

The columns in test.result are the following:

p_value The p-value for each enrichment.

metapone

n_significant m	etabolites
	The number of weighted significant metabolites associated with the enrichment.
n_mapped_metabo	olites
	The number of weighted metabolites associated with the enrichment.
n_metabolites significant met	The number of metabolites associated with the enrichment. abolites
-	A string with the names of significant metabolites that drive the enrichment.
mapped_metabol:	ites
	A string with the names of metabolites that drive the enrichment.
lfdr	The local fdr value for each enrichment.
adjust.p	The enrichment BH-adjusted p-value for each enrichment.
ES	The enrichment score (Avaliable in GSEA test).
NES	The enrichment score normalized to mean enrichment of random samples of the same size (Avaliable in GSEA test).
nMoreExtreme	The number of times a random metabolite set had a more extreme enrichment score value (Avaliable in GSEA test).

Author(s)

Tianwei Yu (<yutianwei@cuhk.edu.cn>) Leqi Tian (<leqitian@link.cuhk.edu.cn>)

References

Small Molecule Pathway Database Mummichog

See Also

pa, hmdbCompMZ

```
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone, p.threshold=0.05,
    n.permu=100,fractional.count.power=0.5, max.match.count=10, use.fgsea = TRUE, use.meta = TRUE)
    hist(ptable(r)[,1])
```

```
# Features based GSEA test
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone, p.threshold=0.05,
    n.permu=100,fractional.count.power=0.5, max.match.count=10, use.fgsea = FALSE, use.meta = FALSE)
hist(ptable(r)[,1])</pre>
```

metaponeResult-class Class "metaponeResult"

Description

This class represents the results of pathway testing. The testing result contain two major components: the significant level of each pathway, and the features matched to each pathway.

Objects from the Class

Objects can be created by calls of the form new("metaponeResult", ...).

Slots

- test.result: a dataframe containing p_value, n_significant metabolites, n_mapped_metabolites, n_metabolites, significant metabolites, mapped_metabolite IDs, lfdr and pathway name.
- mapped.features: A list containing n entries, where n is the number of pathways. Each entry is a data frame, containing the features mapped to this pathway. The information include m.z, retention.time, p.value, statistic, HMDB_ID, theoretical m.z, ion.type, fractional counts.

Methods

- ptable signature(object = "metaponeResult"): return the data.frame of test statistics for each pathway, including p_value, n_significant metabolites, n_mapped_metabolites, n_metabolites, significant metabolites, mapped_metabolite IDs lfdr and and pathway name.
- ftable signature(object = "metaponeResult"): Returns a list containing the mapped features
 in each pathway. Each pathway is represented by a data.frame as an item in the list object.
 The dataframe include information of m.z, retention.time, p.value, statistic, HMDB_ID, theoretical m.z, ion.type, fractional counts.

Author(s)

Tianwei Yu

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neg

Example negative mode data from the Metabolome Atlas of the Aging Mouse Brain

Description

The data is generated from the hypocampus data of the Metabolome Atlas of the Aging Mouse Brain (ST001888) dataset. The p-values and test statistics were obtained by contrasting mouse hypocampus metabolome between prime-age mice and aging mice.

Usage

data("neg")

Format

A data frame with 6947 observations on the following 4 variables.

m.z a numeric vector. The mass-to-charge ratio of the features.

retention.time a numeric vector. The retention time of the features.

p.value a numeric vector. The p-values of the features.

statistic a numeric vector. The test statistics of the features.

References

https://www.metabolomicsworkbench.org/data/DRCCMetadata.php?Mode=Study&DataMode=FactorsData&StudyID=ST0

Examples

data(neg)

ра

Pathway-metabolite match file.

Description

mapps pathways with metabolites.

Usage

data("pa")

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Format

A data frame with 5395 observations on the following 5 variables.

database a character vector pathway.name a character vector HMDB.ID a character vector KEGG.ID a character vector

category a character vector

Source

Small Molecule Pathway Database Mummichog

Examples

data(pa)

pos	Example positive mode data from the Metabolome Atlas of the Aging
	Mouse Brain

Description

The data is generated from the hypocampus data of the Metabolome Atlas of the Aging Mouse Brain (ST001888) dataset. The p-values and test statistics were obtained by contrasting mouse hypocampus metabolome between prime-age mice and aging mice.

Usage

data("pos")

Format

A data frame with 10085 observations on the following 4 variables.

m.z a numeric vector. The mass-to-charge ratio of the features.

retention.time a numeric vector. The retention time of the features.

p.value a numeric vector. The p-values of the features.

statistic a numeric vector. The test statistics of the features.

References

https://www.metabolomicsworkbench.org/data/DRCCMetadata.php?Mode=Study&DataMode=FactorsData&StudyID=ST0

Examples

data(pos)

ptable

Description

return the data. frame of test statistics for each pathway.

Usage

```
## S4 method for signature 'metaponeResult'
ptable(object)
```

Arguments

object A metaponeResult object.

Details

Includes p_value, n_significant metabolites, n_mapped_metabolites, n_metabolites, significant metabolites, mapped_metabolite IDs and pathway name.

Value

The method returns a data frame with 6 columns: "p_value", "n_significant metabolites", "n_mapped_metabolites", "n_metabolites", "significant metabolites", "mapped_metabolites".

Author(s)

Tianwei Yu <yutianwei@cuhk.edu.cn>

See Also

ftable

```
data(hmdbCompMZ.metapone)
data(pa)
data(pos)
data(neg)
dat <- list(pos, neg)
type <- list("pos", "neg")
r<-metapone(dat, type, pa, hmdbCompMZ=hmdbCompMZ.metapone,
    p.threshold=0.05,n.permu=100,fractional.count.power=0.5, max.match.count=10)
head(ptable(r))</pre>
```

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