

Package ‘omicReposome’

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Title Exposome and omic data association and integration analysis

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Description omicReposome systematizes the association evaluation between exposures and omic data, taking advantage of MultiDataSet for coordinated data management, reposome for exposome data definition and limma for association testing. Also to perform data integration mixing exposome and omic data using multi co-inherent analysis (omicade4) and multi-canonical correlation analysis (PMA).

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add_cls	<i>Method to add an ExosomeClust to a MultiDataSet</i>
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Description

This method allows to insert an object of class [ExosomeClust](#) as an independent dataset into an object of class [MultiDataSet](#).

Usage

```
add_cls(object, clsSet, ...)

## S4 method for signature 'MultiDataSet,ExosomeClust'
add_cls(object, clsSet, ...)
```

Arguments

- object An object of class [MultiDataSet](#).
- clsSet An object of class [ExosomeClust](#).
- ... Arguments given to [add_eset](#) from [MultiDataSet](#).

Value

A [MultiDataSet](#) with the [ExpressionSet](#) added as an independent dataset.

Examples

```
data("eclust", package = "rexposome")
library(MultiDataSet)
md <- new("MultiDataSet")
names(md)
md <- add_cls(md, expo_c)
names(md)
```

add_exp

Method to add an ExposomeSet to a MultiDataSet

Description

This method allows to insert an object of class [ExposomeSet](#) as an independent dataset into an object of class [MultiDataSet](#).

Usage

```
add_exp(object, expoSet, warnings = TRUE, ...)
## S4 method for signature 'MultiDataSet,ExposomeSet'
add_exp(object, expoSet, warnings = TRUE,
        ...)
```

Arguments

object	An object of class MultiDataSet .
expoSet	An object of class ExposomeSet .
warnings	(default TRUE) If set to FALSE warnings will not be displayed.
...	Arguments given to add_eset from MultiDataSet .

Value

A [MultiDataSet](#) with the [ExpressionSet](#) added as an independent dataset.

Examples

```
data("exposome", package = "rexposome")
library(MultiDataSet)
md <- new("MultiDataSet")
names(md)
md <- add_exp(md, expo)
names(md)
```

asr	<i>ResultSet for testing and illustration purposes</i>
-----	--

Description

ResultSet created using **association** method, testing proteome association to exposome ("mds"), adjusted by sex and age.

Usage

```
data("asr")
```

Format

An object of class **ResultSet** of length 15.

Value

A **ResultSet** object.

Examples

```
data("asr", package = "omicReXposome")
asr
```

association	<i>#</i>	<i>Method to perform an association study between transcriptome and exposome</i>
-------------	----------	--

Description

This function allows to perform an association study between gene expression from microarray and the exposome. An ExpressionSet is the object storing the gene expression and an ExposomeSet the one storing the exposome. Both of them need to be encapsulated in a MultiDataSet. The association study is performed through standard limma pipeline. The function allows to perform multiple tests using the argument exposures.

Usage

```
association(object, formula, expset, omicset, set = "exposures",
method = "ls", ..., baselevels, sva = "none", vfilter = NULL,
verbose = FALSE, warnings = TRUE)

## S4 method for signature 'MultiDataSet'
association(object, formula, expset, omicset,
set = "exposures", method = "ls", ..., baselevels, sva = "none",
vfilter = NULL, verbose = FALSE, warnings = TRUE)
```

Arguments

<code>object</code>	A <code>MultiDataSet</code> object containing at last one omic data-sets like <code>ExpressionSet</code> , <code>MethylationSet</code> ... and, at last, one <code>ExposomeSet</code> .
<code>formula</code>	formula to be evaluated by each exposure (or phenotype, see <code>set</code> argument). It should not contain any exposures (or phenotype), it will be added automatically when evaluated.
<code>expset</code>	Name of the <code>ExposomeSet</code> in <code>object</code> .
<code>omicset</code>	Name of the omic data-set in <code>object</code> .
<code>set</code>	(default "exposures") Can take value "exposures" to test the association of the exposures in the <code>ExposomeSet</code> vs. the features in the omic data-set. If takes "phenotypes" all phenotypes in <code>ExposomeSet</code> are tested.
<code>method</code>	(default "lm") Check <code>limma</code> help pages.
...	Arguments passed to <code>limma</code> 's <code>lmFit</code> .
<code>baselevels</code>	(optional) If set, must be a labeled vector with the default base level for categorical exposures.
<code>sva</code>	(default "none"). This argument can take value "none" to do not apply SVA. Value "fast" will run SVA using <code>isva</code> and <code>SmartSVA</code> . Value "slow" will run SVA using <code>sva</code> .
<code>vfilter</code>	(default NULL). Only used when <code>sva = "slow"</code> . Numeric number of probes used in <code>sva</code> . Recomended ~10% of real probes.
<code>verbose</code>	(default FALSE) If set to TRUE, a series of messages describing the process are shown.
<code>warnings</code>	(default TRUE) If set to TRUE, a series of warnings are shown when required user attention.

Value

An object of class `ResultSet`.

Examples

```
library(MultiDataSet)
data(brge_prot, package = "brgedata")
data(brge_expo, package = "brgedata")
mds <- createMultiDataSet()
mds <- add_eset(mds, brge_prot, dataset.type = "proteines")
mds <- add_eset(mds, brge_expo, dataset.type = "exposures", GRanges = NA)

asr <- association(mds, formula = Asthma ~ Sex + Age,
                    expset = "exposures", omicset = "proteines")
asr
```

crossomics*Function to perform a Transcriptome-Wide Association Study*

Description

This function allows to perform a Transcriptome-Wide Association Study by using an ExposomeSet and an ExpressionSet. It allows to perform an adjustment using Surrogate Variable Analysis (from R package sva).

Usage

```
crossomics(object, method = "mcca", ncomponents = 2, ..., na.rm = FALSE,
           permute = c(100, 3), verbose = FALSE, warnings = TRUE)

## S4 method for signature 'MultiDataSet'
crossomics(object, method = "mcca",
           ncomponents = 2, ..., na.rm = FALSE, permute = c(100, 3),
           verbose = FALSE, warnings = TRUE)
```

Arguments

object	A MultiDataSet object containing at least two data-sets like ExposomeSet, ExpressionSet, MethylationSet...
method	(default "mcca") It can take values "mcca" for Multiple Canonical Correlation Analysis or "mcia" for Multiple Co-Inertia Analysis.
ncomponents	(default 2) Number of components to be estimated.
...	Other arguments given to mcia (from omicade4) or to MultiCCA (from PMA).
na.rm	(default FALSE) If method was set to "mcca" and na.rm was set to TRUE, features containing missing values are removed.
permute	(default c(100, 3)). If method="mcca" and this argument is set to NULL no permutation test to tune-up the parameters for MultiCCA. When filled, permute[1] corresponds to the number of permutations (default in MultiCCA.permute is 25) and permute[2] the number of iterations (default in MultiCCA.permute is 3).
verbose	(default FALSE) If set to TRUE, a series of messages describing the process are shown.
warnings	(default TRUE) If set to TRUE, a series of warnings are shown when required user attention.

Value

An object of class [ResultSet](#).

Examples

```
library(MultiDataSet)
library(rexposome)
data(brge_prot, package = "brgedata")
data(brge_expo, package = "brgedata")
mds <- createMultiDataSet()
mds <- add_eset(mds, brge_prot, dataset.type = "proteines")
mds <- add_eset(mds, imputation(brge_expo),
                dataset.type = "exposures", GRanges = NA)

crs <- crossomics(mds, method = "mcia")
crs
```

crs

ResultSet *for testing and illustration purposes*

Description

ResultSet created using `crossomics` method, selecting "mcia" method. Result from the integration of proteome and exposome data ("mds").

Usage

```
data("crs")
```

Format

An object of class ResultSet of length 1.

Value

A ResultSet object.

Examples

```
data("crs", package = "omicReXposome")
crs
```

<code>getIntegration</code>	<i>Method to extract integration-feature result from a ResultSet</i>
-----------------------------	--

Description

Homologous methods from MultiDataSet (getAssociation) but for ResultSet created by [crossomics](#). It Returns a data.frame with the result from mcia (omicade4) or from MultiCCA (PMA).

Usage

```
getIntegration(object, ...)

## S4 method for signature 'ResultSet'
getIntegration(object, ...)
```

Arguments

<code>object</code>	An object of class ResultSet obtained from
...	NOT USED

Value

A data.frame

Examples

```
data("crs", package = "omicRexposome")
class(getIntegration(crs))
```

<code>mds</code>	<i>MultiDataSet for testing and illustration purposes</i>
------------------	---

Description

MultiDataSet containing both proteome data-set and exposome data-set.

Usage

```
data("mds")
```

Format

An object of class MultiDataSet of length 2.

Value

A MultiDataSet object.

Examples

```
data("mds", package = "omicRexposome")
mds
```

omicRexposome

omicRexposome: Package for exposome and omic data associatin and integration

Description

omicRexposome: Package for exposome and omic data associatin and integration

exposome-omic data association study

The packages offers the function [association](#) that allows to perform an association study using transcriptome, methylome, etc. as dependent variable and exposome data as independent variable. The function relies on limma pipeline and generates an object of class `ResultSet`, that can be plotted using [plotAssociation](#).

exposome-omic data integration study

The packages offers the function [crossomics](#) that allows to perform two types of integration study: Multi Canonical Correlation Analysis and Multi Co-Inertia Analysis. The function allows to use any type and number of datasets (aka. exposome transcriptome, methylome, etc.). The function generates an object of class `ResultSet`, that can be plotted using [plotIntegration](#).

plotAssociation

Function to draw de result of an association study

Description

This function draws two type of plots for the `ResultSet` from association functions

Usage

```
plotAssociation(object, rid = 1, coef = 2, contrast = 1,
                type = c("manhattan", "qq", "volcano"), tPV = NULL, tFC = NULL,
                show.effect = FALSE)

## S4 method for signature 'ResultSet'
plotAssociation(object, rid = 1, coef = 2,
                contrast = NULL, type = c("manhattan", "qq", "volcano"), tPV = NULL,
                tFC = NULL, show.effect = FALSE)
```

Arguments

<code>object</code>	An object of class ResultSet obtained from <code>assoc_*</code> functions.
<code>rid</code>	(default 1) Index or name of the test to be plotted.
<code>coef</code>	(default 2) Index of the coefficient to be extracted.
<code>contrast</code>	(default 1) When code corresponds to a multicategorical variable, <code>contasr</code> selects the comparison.
<code>type</code>	Can take "volcano", "qq", "manhattan" and "protein". "protein" lot is a type of Manhattan plot designed for protein association analysis.
<code>tPV</code>	(optional) Threshold for P.Value when <code>type="volcano"</code> .
<code>tFC</code>	(optional) Threshold for Fold Change or Effect when <code>type="volcano"</code> .
<code>show.effect</code>	(default FALSE) If set to TRUE, when <code>type="volcano"</code> the X-axis will show $2^{\log FC}$ instead of $\log FC$.

Value

A ggplot2 object

See Also

[plotIntegration](#) for plotting integration results. [association](#) to create a `ResultSet` to be passed to this function.

Examples

```
data("asr", package = "omicRexposome")
plotAssociation(asr, type = "qq")
plotAssociation(asr, type = "volcano")
```

plotHits

Plot number of hits per result in ResultSet

Description

This method draws a barplot with the number of hits in each result stored in the given [ResultSet](#).

Usage

```
plotHits(object, th = 0.05, width = 0.75)

## S4 method for signature 'ResultSet'
plotHits(object, th = 0.05, width = 0.75)
```

Arguments

object	An object of class ResultSet
th	(default 0.05) Threshold (p-value) to consider a result as a hit.
width	(default 0.70) width of the bar

Value

A ggplot2 object

See Also

[plotLambda](#) for a graphical representation of the lambda score per analysis, [tableLambda](#) for the lambda score per analysis, [tableHits](#) for the histograms per analysis

Examples

```
data(asr, package = "omicReposome")
plotHits(asr)
```

plotIntegration *Function to draw the result of an integration study*

Description

This function draws a plot for the ResultSet from integration function

Usage

```
plotIntegration(object, cmpX = 1, cmpY = 2, lb.th = 0.2,
               legend.show = TRUE, colors, ...)

## S4 method for signature 'ResultSet'
plotIntegration(object, cmpX = 1, cmpY = 2,
               lb.th = 0.2, legend.show = TRUE, colors, ...)
```

Arguments

object	An object of class ResultSet obtained from crossomics .
cmpX	(default 1) Value of the X-axis when plotting results from mcia .
cmpY	(default 2) Value of the Y-axis when plotting results from mcia .
lb.th	(default 0.20) Threshold to place labels on radar chart drawn when plotting results from MultiCCA .
legend.show	(default TRUE) If set to FALSE, right legend of radar plot is hidden when plotting results from MultiCCA .
colors	(optional) Names vector with the colors used to draw each dataset. Used when plotting results from MultiCCA . If missing, random colors are chosen.
...	Optional arguments are given to plot from omicade4 package (argument axes is filled with values from cmpX and cmpY).

Value

A ggplot2 object

See Also

[plotAssociation](#) for plotting association results. [crossomics](#) to create a **ResultSet** to be passed to this function.

Examples

```
data("crs", package = "omicReposome")
plotIntegration(crs)
```

plotLambda

Plot lambda score for all results in a ResultSet

Description

This method draws a baplor with the lambda score of each result in the given [ResultSet](#).

Usage

```
plotLambda(object, width = 0.75)

## S4 method for signature 'ResultSet'
plotLambda(object, width = 0.75)
```

Arguments

object	An object of class ResultSet
width	(default 0.70) width of the bar

Value

A ggplot2 object

See Also

[plotHits](#) for a graphical representation of the hits per analysys, [tableLambda](#) for the lambda score per analysys, [tableHits](#) for the hists per analysys

Examples

```
data("asr", package = "omicReposome")
plotLambda(asr)
```

snpToContinuous	<i>Transforms the discrete genotype from a snpSet to a matrix of a continuous variable.</i>
-----------------	---

Description

The function converts the categorical variable of SNPs to a continuous variable by normalizing each SNP as described in Abraham G. and Inouye M. 2014 (DOI: 10.1371/journal.pone.0093766).

Usage

```
snpToContinuous(snpSet, verbose = FALSE)
```

Arguments

snpSet	An object of class <code>snpSet</code> with set calls slot .
verbose	If set to TRUE, messages will be shown.

Value

An matrix of the calls of the SNPs converted to a continuous variable.

See Also

[crossomics](#) use this function

tableHits	<i>Counts the number of hits on the results stored in a ResultSet</i>
-----------	---

Description

Given a threshold it counts the number of hits in each result in the given [ResultSet](#).

Usage

```
tableHits(object, th = 0.05)

## S4 method for signature 'ResultSet'
tableHits(object, th = 0.05)
```

Arguments

object	An object of class ResultSet
th	(default 0.05) Threshold (p-value) to considere a result as a hit.

Value

A labeled numeric vector with the exposures and the number of hits.

See Also

[tableLambda](#) for the lambda score per analysys, [plotLambda](#) for a graphical representation of the lambda score per analysys, [plotHits](#) for a graphical representation of the hists per analysys

Examples

```
data("asr", package = "omicReposome")
tableHits(asr)
```

tableLambda

Compute a lambda score on the results stored in a ResultSet

Description

Compute lambda score on each result in the given [ResultSet](#) by using [lambdaClayton](#).

Usage

```
tableLambda(object, trim = 0.5)

## S4 method for signature 'ResultSet'
tableLambda(object, trim = 0.5)
```

Arguments

object	An object of class ResultSet
trim	(default 0.5) percentage of right omitted values for lambdaClayton .

Value

Returns a `data.frame` having the exposures and the computed lambda score.

A labeled numeric vector with the lambda score for each exposure.

See Also

[tableHits](#) for the number of hits per analysys, [plotHits](#) for a graphical representation of the hists per analysys, [plotLambda](#) for a graphical representation of the lambda score per analysys

Examples

```
data("asr", package = "omicReposome")
tableLambda(asr)
```

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