Package 'MetaPcor'

February 28, 2023

Type Package

Usage

Hite Meta-analysis of gene expression using partial correlation as effect size
Version 0.1.0
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Description This package performs meta-analysis of gene expression studies using partial correlation as effect size, to provide the means to construct a causal association gene network
License GPL-3
Encoding UTF-8
NeedsCompilation no
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DEG_meta
Description Differential expression analysis, which keeps only the statistically significant genes from a give
set of gene expression studies

DEG_meta(folder_path, cases, controls)

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Arguments

folder_path A folder path where the gene expression studies are stored cases How the cases are named in the gene expression studies controls How the controls are named in the gene expression studies

Value

A list of data.tables containing gene expression columns of the statistically significant genes that occured from the differential epxression analysis

Author(s)

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Examples

```
DEG_ES <- DEG_meta(folder_path = 'studies/', cases = 'CASE',
controls = 'CONTROL')</pre>
```

enirchment_analysis

Description

A function to conduct enrichment analysis with the statistically significant genes that occured from the meta-analysis

Usage

```
enirchment_analysis(x)
```

Arguments

Х

A dataframe object containing the results of the meta_pcor function

Value

A list with the results of the enrichment analysis with the gProfiler tool ([1]: A dataframe with the results of the enrichment and [2] an interactive Manhattan plot with the statistically significant terms from the sources)

Author(s)

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References

Uku Raudvere, Liis Kolberg, Ivan Kuzmin, Tambet Arak, Priit Adler, Hedi Peterson, Jaak Vilo, g:Profiler: a web server for functional enrichment analysis and conversions of gene lists (2019 update), Nucleic Acids Research, Volume 47, Issue W1, 02 July 2019, Pages W191–W198, https://doi.org/10.1093/nar/gkz3

See Also

https://biit.cs.ut.ee/gprofiler/gost

Examples

```
ea_results <- enrichment_analysis(meta_pcor_results)</pre>
```

meta_pcor

Partial Correlation Meta-Analysis Function

Description

This function performs meta-analysis of partial correlation coefficients using neighborhood selection approach or shrinkage methods. The function accepts data from multiple studies as input and outputs a meta-analysis result that represents a summary of the partial correlation coefficients obtained from the individual studies.

Usage

```
meta_pcor(file_names, option, method, meta_method= "random",
pvalue_thres = NULL, fdr_thres = NULL,
coef_thres = NULL,11 = NULL ,12 = NULL, norm_data = NULL, norm_method = NULL)
```

Arguments

_	
file_names	a character vector containing the file names of the data to be analyzed.
option	The meta-analysis option to run (integer). 1: Pearson correlation and partial correlation meta-analysis 2: Partial correlation meta-analysis with thresholds 3: Partial correlation meta-analysis without thresholds 4: Differential expression and partial correlation meta-analysis
method	a character string indicating the method to be used to estimate partial correlation coefficients. Available options are "sparse" and "shrinkage".
meta_method	a character string indicating the method to be used for the meta-analysis of the partial correlation coefficients. Available options are "random" and "fixed". Default is "random".
<pre>pvalue_thres</pre>	a numeric value specifying the threshold for p-values.
fdr_thres	a numeric value specifying the threshold for false discovery rate (FDR).
coef_thres	a numeric value specifying the threshold for correlation coefficient magnitude.
11	a numeric value specifying the parameter for neighborhood selection approach.
12	a numeric value specifying the parameter for neighborhood selection approach.
norm_data	a character string indicating whether data normalization should be performed. Available options are "YES" and "NO". Default is "NO".
norm_method	a character string indicating the method to be used for normalization. Available

options are "RPKM" and "Z_SCOR_STAND", "LOG2" and "QUANT_NORM".

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Value

A data.table object is returned, containing the results of the meta-analysis.

Author(s)

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References

Uku Raudvere, Liis Kolberg, Ivan Kuzmin, Tambet Arak, Priit Adler, Hedi Peterson, Jaak Vilo, g:Profiler: a web server for functional enrichment analysis and conversions of gene lists (2019 update), Nucleic Acids Research, Volume 47, Issue W1, 02 July 2019, Pages W191–W198, https://doi.org/10.1093/nar/gkz3

See Also

https://biit.cs.ut.ee/gprofiler/gost

Examples

```
metapcor_result <- meta_pcor(GEO_names=c("GSE76427"),
target_namespace = c('ILLUMINA_HUMANHT_12_V4'), option = 2,
method = "sparse", meta_method = "random",
pvalue_thres = 0.01, l1 = 0.8, l2 = 0)

metapcor_result <- meta_pcor(folder_path = "/path/to/data/folder",
option = 1, method = "sparse",meta_method = "random",l1 = 0.8, l2 = 0)</pre>
```

my_meta

Description

A function that performs meta-analysis of partial correlations

Usage

```
my_meta(correlations, method)
```

Arguments

correlations A partial correlation data.table object, which occurs from the results of the par-

tial_neighborhood or the partial_shrinkage functions

method Meta-analysis method to calculate the effect sizes ('random' or 'fixed')

Value

A data.table containing the results of the meta-analysis

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Examples

```
my_meta_results <- my_meta(correlations = sparse_partial_corr, method = 'random')</pre>
```

Description

network_plot

A function which plots an interactive causal association gene netowrk with the GraphViz package

Usage

```
network_plot(x)
```

Arguments

х

A data.table object which contains the results of the meta_pcor function

Value

An interactive network plot of the causal association gene network

Author(s)

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Examples

```
# Network plot
network_plot(meta_pcor_results)

pcor_neighborhood
```

Description

A function to calculate sparse partial correlation estimation with neighborhood selection approach method for each study

Usage

```
pcor_neighborhood(list_of_files, 11, 12=0, significant, pvalue_thres=NULL,
fdr_thres= NULL, coef_thres = NULL)
```

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Arguments

list_of_files	A list of files that contain the transposed gene expression studies as data.table objects
11	11 Norm penalty. $11 = 0.8$ is suggested (float)
12	12 value for sparse partial correlation calculation. $12 = 0.0$ is suggested (float)
significant	A boolean option ("TRUE" or "FALSE") to choose only the statistically significant pairs of partial correlations, either with a p-value or a FDR or a coefficient threshold
pvalue_thres	p-value threshold for the meta-analysis (float)
fdr_thres	FDR value threshold for the meta-analysis (float)
coef_thres	Coefficient threshold for the meta-analysis (float)

Value

A data.table object is returned with the partial correlations pairs

Author(s)

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See Also

https://github.com/cran/space

Examples

```
# Calculate sparse partial correlation estimation with neighborhood selection approach method for each study
sparse_partial_corr <- pcor_neighborhood(list_of_files = list_of_studies,
11 = 0.8, 12 = 0.0, significant = TRUE, pvalue_thres = 0.05)</pre>
```

pcor_shrinkage

Description

A function to calculate sparse partial correlation estimation with the shrinkage method for each study.

Usage

```
pcor_shrinkage(x)
```

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Arguments

list_of_files A list of files that contain the transposed gene expression studies as data.table

objects

significant A boolean option ("TRUE" or "FALSE") to choose only the statistically signifi-

cant pairs of partial correlations, either with a p-value or a FDR or a coefficient

threshold

pvalue_thres p-value threshold for the meta-analysis (float)

fdr_thres FDR value threshold for the meta-analysis (float)

coef_thres Coefficient threshold for the meta-analysis (float)

Value

A data.table object is returned with the partial correlations pairs

Author(s)

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Examples

```
shrinkage_partial_corr <- pcor_shrinkage(list_of_files, significant = TRUE, pvalue_thres = 0.05)</pre>
```

volc_plot

Description

A function which produces an interactive volcano plot with the Plotly package

Usage

```
volc_plot_plotly(x, pval_thres, coeff_thres)
```

Arguments

x A data.table object which contains the results of the meta_pcor function

pval_thres A p-value threshold for the volcano (float)

coeff_thres A coefficient threshold for the volcano plot (float)

Value

An interactive volcano plot

Author(s)

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Examples

```
# Volcano Plot
volc_plot<- volc_plot_plotly(meta_pcor_results, pval_thres = 0.05,
coeff_thres = 0.2)</pre>
```

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