Package 'MetaPhOR'

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Type Package

Title Metabolic Pathway Analysis of RNA

Version 1.2.0

Description MetaPhOR was developed to enable users to assess metabolic dysregulation using transcriptomic-level data (RNA-sequencing and Microarray data) and produce publicationquality figures. A list of differentially expressed genes (DEGs), which includes fold change and p value, from DESeq2 or limma, can be used as input, with sample size for MetaPhOR, and will produce a data frame of scores for each KEGG pathway. These scores represent the magnitude and direction of transcriptional change within the pathway, along with estimated pvalues.MetaPhOR then uses these scores to visualize metabolic profiles within and between samples through a variety of mechanisms, including: bubble plots, heatmaps, and pathway models. License Artistic-2.0 **Encoding** UTF-8 RoxygenNote 7.2.1 Imports utils, ggplot2, ggrepel, stringr, pheatmap, grDevices, stats, clusterProfiler, RecordLinkage, RCy3 **Depends** R (>= 4.2.0) biocViews Metabolomics, RNASeq, Pathways, GeneExpression, DifferentialExpression, KEGG, Sequencing, Microarray Suggests BiocStyle, knitr, rmarkdown, kableExtra VignetteBuilder knitr LazyData false SystemRequirements Cytoscape (>= 3.9.0) for the cytoPath() examples git url https://git.bioconductor.org/packages/MetaPhOR git branch RELEASE 3 17 git_last_commit 9a86f8e git last commit date 2023-04-25 Date/Publication 2023-10-15 Author Emily Isenhart [aut, cre], Spencer Rosario [aut]

Maintainer Emily Isenhart <emily.isenhart@roswellpark.org>

R topics documented:

bubblePlot																	•	•	•	•						2
cytoPath																										3
datasummary																										4
metaHeatmap																										4
pathwayAnalys	is			•		•		•	•			•					•	•			•	•	•	•		5
pathwayList .		•				•		•	•			•		•		•	•	•		•	•	•	•	•	•	6
																										_
																										7

Index

bubblePlot

Create a Bubble Plot for Individual Samples

Description

Create a Bubble Plot for Individual Samples

Usage

bubblePlot(scorelist, labeltext, labelsize = 0.25)

Arguments

scorelist	dataframe(1) the output of Pathway Analysis fun
labeltext	character(1) what to label points by: LogFC or Pval
labelsize	numeric(1) size of text labels for points

Value

bubblePlot() returns a bubble plot using pathway scores, pval, logfc

Examples

cytoPath

Description

requires the package RCy3 and a local instance of Cytoscape

Usage

```
cytoPath(
    pathway,
    DEGpath,
    figpath,
    genename,
    headers = c("log2FoldChange", "padj")
)
```

Arguments

pathway	character, the name of the pathway to be visualized
DEGpath	character, the path to a DEG file by DESeq2 or limma
figpath	character, the path to which the figure will be saved
genename	character, column name with HUGO Gene Names in DEG file
headers	character vector of length 2 in the form c(log fold change col name, adjusted p value col name)

Value

cytoPath() Returns a Cytoscape figure of DEG data on rWikiPathways

Examples

```
cytoPath(pathway = "Tryptophan Metabolism",
    DEGpath = system.file("extdata/BRCA_DEGS.csv", package = "MetaPhOR"),
    figpath = file.path(tempdir(), "example_map"),
    genename = "X",
    headers = c("logFC", "adj.P.Val"))
```

```
datasummary
```

Description

MetaPhOR was developed to enable users to assess metabolic dysregulation using transcriptomiclevel data (RNA-sequencing and Microarray data) and produce publication-quality figures. A list of differentially expressed genes (DEGs), which includes fold change and p value, from DESeq2 or limma, can be used as input, with sample size for MetaPhOR, and will produce a data frame of scores for each KEGG pathway. These scores represent the magnitude and direction of transcriptional change within the pathway, along with estimated p-values. MetaPhOR then uses these scores to visualize metabolic profiles within and between samples through a variety of mechanisms, including: bubble plots, heatmaps, and pathway models.

Author(s)

Maintainer: Emily Isenhart <emily.isenhart@roswellpark.org>

Authors:

· Spencer Rosario

metaHeatmap

Create a Heatmap for Comparing Multiple Samples

Description

Create a Heatmap for Comparing Multiple Samples

Usage

```
metaHeatmap(scorelist, samplenames, pvalcut = 0.05)
```

Arguments

scorelist	list of outputs from pathwayAnalysis()
samplenames	vector of samples names for axis labels
pvalcut	numeric, the p val over which pathways will not be included

Value

metaHeatmap() returns a heatmap of significant dysregulated pathways for each sample included

pathwayAnalysis

Examples

pathwayAnalysis Metabolic Pathway Analysis of RNAseq Data

Description

Metabolic Pathway Analysis of RNAseq Data

Usage

```
pathwayAnalysis(
    DEGpath,
    genename,
    sampsize,
    iters = 1e+05,
    headers = c("log2FoldChange", "padj")
)
```

Arguments

DEGpath	character, the path to a txt or csv DEG file
genename	character, column name with HUGO Gene Names in DEG file
sampsize	numeric, the sample size of the experiment to be analyzed
iters	numeric, the number of iterations of resampling to perform in bootstrapping
headers	character vector of length2 in the form c(log fold change col name, adjusted p value col name)

Value

pathwayAnalysis() returns a dataframe of pathway scores and pvals

Examples

```
#iterations (iters) of resampling in bootstraping set to 30,000 for speed
#100,000 iterations recommended for improved power
set.seed(1234)
scores <- pathwayAnalysis(</pre>
```

scores

```
pathwayList List Available Metabolic rWikiPathways
```

Description

List Available Metabolic rWikiPathways

Usage

pathwayList()

Value

pathwayList() returns a list of rWikiPathways for use in CytoPath()

Examples

pathwayList()

Index

bubblePlot, 2

cytoPath, 3

datasummary,4

metaHeatmap, 4
MetaPhOR (datasummary), 4
MetaPhOR-package (datasummary), 4

pathwayAnalysis,5
pathwayList,6