

Package ‘EnrichIntersect’

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Type Package

Title Enrichment Analysis and Intersecting Sankey Diagram

Version 1.0

URL <https://github.com/ocbe-uio/EnrichIntersect>

BugReports <https://github.com/ocbe-uio/EnrichIntersect/issues>

Description

A flexible tool for enrichment analysis based on user-defined sets. It allows users to perform over-representation analysis of the custom sets among any specified ranked feature list, hence making enrichment analysis applicable to various types of data from different scientific fields. 'EnrichIntersect' also enables an interactive means to visualize identified associations based on, for example, the mix-lasso model (Zhao et al., 2022 <[doi:10.1016/j.isci.2022.104767](https://doi.org/10.1016/j.isci.2022.104767)>) or similar methods.

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Depends R (>= 4.0)

Encoding UTF-8

LinkingTo Rcpp, RcppArmadillo

Imports Rcpp, ggplot2, dplyr, networkD3, jsonlite, htmlwidgets, webshot2, stats, methods, grDevices

Suggests knitr

LazyData true

VignetteBuilder knitr

NeedsCompilation yes

Config/roxygen2/version 8.0.0

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Contents

cancers_drug_groups	2
cancers_genes_drugs	3
enrichment	3
intersectSankey	4
Index	6

cancers_drug_groups *Data set 'cancers_drug_groups'*

Description

Data set 'cancers_drug_groups' is a list including a score dataframe with 147 drugs as rows and 19 cancer types as columns, and a dataframe with 9 self-defined drug groups (1st column) of the 147 drugs (2nd column). More details of the data can be found in the results of paper 'Tissue-specific identification of multi-omics features for pan-cancer drug response prediction' by Zhao Z., Wang S, Zucknick M. and Aittokallio T. (2022).

Usage

```
cancers_drug_groups
```

Format

An object of class `list` of length 2.

Value

No return value.

References

Zhao Z, Wang S, Zucknick M and Aittokallio T (2022). *Tissue-specific identification of multi-omics features for pan-cancer drug response prediction*. *iScience*, 25(8): 104767.

Examples

```
# Load data
data(cancers_drug_groups, package = "EnrichIntersect")
```

cancers_genes_drugs	<i>Data set 'cancers_genes_drugs'</i>
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Description

Data set 'cancers_genes_drugs' is an array with association scores between 56 genes (1st dimension), three cancer types (2nd dimension) and two drugs (3rd dimension). More details of the data can be found in the results of paper 'Tissue-specific identification of multi-omics features for pan-cancer drug response prediction' by Zhao Z., Wang S. Zucknick M. and Aittokallio T. (2022).

Usage

```
cancers_genes_drugs
```

Format

An object of class array of dimension 56 x 3 x 2.

Value

No return value.

References

Zhao Z, Wang S, Zucknick M and Aittokallio T (2022). *Tissue-specific identification of multi-omics features for pan-cancer drug response prediction*. *iScience*, 25(8): 104767.

Examples

```
# Load data
data(cancers_genes_drugs, package = "EnrichIntersect")
```

enrichment	<i>Plot enrichment map</i>
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Description

Plot enrichment map through a vector or matrix of scores and a self-defined set that summarizes a few groups of the names of the vector or matrix.

Usage

```
enrichment(
  x,
  custom.set,
  alpha = 0,
  normalize = TRUE,
  permute.n = 100,
  padj.method = "none",
  pvalue.cutoff = 0.05,
  angle = 45,
  match.feature = c("rownames", "colnames"),
  ...
)
```

Arguments

x	a vector or matrix of scores to be enriched
custom.set	a self-defined set. The first column contains feature names, and the second column, preferably named "group", contains group names.
alpha	exponent weight of the score of ordered features.
normalize	logical value to determine if normalizing enrichment scores.
permute.n	number of custom-set permutations for significance testing.
padj.method	correction method passed to stats::p.adjust.
pvalue.cutoff	cutoff for both unadjusted and adjusted p-value.
angle	angle of rotating x-axis labels.
match.feature	one of "rownames" or "colnames". Default is "rownames", which keeps the original EnrichIntersect behavior.
...	other arguments

Value

A list including S, pvalue, and g.

 intersectSankey

Plot Sankey diagram for intersecting sets

Description

Plot Sankey diagram for intersecting set through an array

Usage

```
intersectSankey(
  x,
  out.fig = NULL,
  color = NULL,
  step.names = c("Levels", "Variables", "Tasks"),
  fontSize = c(20, 13, 20),
  nodePadding = 11,
  nodeWidth = 5,
  margin = list(right = 180),
  ...
)
```

Arguments

x	an array for constructing intersecting set
out.fig	print the figure into "html", "pdf" or "png" file. Default is NULL with R graphics device
color	a vector of colors corresponding to individual tasks
step.names	names of the three dimensions of the array x, i.e. names of multiple levels, intermediate variables and tasks. Default is c("Levels", "Variables", "Tasks"). If step.names=NULL, it will not show the names
fontSize	a value or vector of three values. If it is one value, it is the font size for all labels. But a vector of three values specifies the font size of the labels in the left, middle and right, respectively. Default is c(20, 10, 20)
nodePadding	numeric essentially influences the width height
nodeWidth	numeric width of each node
margin	an integer or a named list/vector of integers for the plot margins
...	graphics parameters to be passed to sankeyNetwork() from R package networkD3

Value

An object of a D3 JavaScript intersecting Sankey diagram for visualising associations based on the input array.

Examples

```
# Data set 'cancers_genes_drugs' is an array with association scores between 56 genes (1st
# dimension), three cancer types (2nd dimension) and two drugs (3rd dimension)
data(cancers_genes_drugs, package = "EnrichIntersect")
```

```
intersectSankey(cancers_genes_drugs, step.names = c("Cancers", "Genes", "Drugs"))
```

Index

* datasets

cancers_drug_groups, [2](#)

cancers_genes_drugs, [3](#)

cancers_drug_groups, [2](#)

cancers_genes_drugs, [3](#)

enrichment, [3](#)

intersectSankey, [4](#)