

Package ‘DrugSim2DR’

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Type Package

Title Predict Drug Functional Similarity to Drug Repurposing

Version 0.1.0

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Description A systematic biology tool was developed to repurpose drugs via a drug-drug functional similarity network. 'DrugSim2DR' first predict drug-drug functional similarity in the context of specific disease, and then using the similarity constructed a weighted drug similarity network. Finally, it used a network propagation algorithm on the network to identify drugs with significant target abnormalities as candidate drugs.

License GPL (>= 2)

Encoding UTF-8

LazyData true

RoxygenNote 7.2.1

Imports igraph,
stats,
pheatmap,
CheminR,
rvest,
base,
sp,
tidyR,
reshape2

Suggests knitr,
rmarkdown

VignetteBuilder knitr

Depends R (>= 3.6)

R topics documented:

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| | |
|------------|-------------------|
| CalDEscore | <i>CalDEscore</i> |
|------------|-------------------|

Description

Function "CalDEscore" uses gene expression to calculate differential expression level.

Usage

```
CalDEscore(exp, Label)
```

Arguments

| | |
|-------|--|
| exp | A gene expression profile of interest (rows are genes, columns are samples). |
| Label | A character vector consist of "0" and "1" which represent sample class in the gene expression profile. "0" means normal sample and "1" means disease sample. |

Value

A matrix with one column of zscore.

Examples

```
# Obtain the example data
GEP<-Gettest("GEP")
label<-Gettest("label")
# Run the function
DEscore<-CalDEscore(GEP,label)
```

| | |
|-------------|---|
| datasummary | <i>datasummary: Custom Data Summaries</i> |
|-------------|---|

Description

Easily generate custom data frame summaries

Author(s)

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*DrugReposition**DrugReposition*

Description

The function "DrugReposition" is used in drug repositioning by calculating the eigenvector centrality of drugs.

Usage

```
DrugReposition(DE,nperm = 100,r = 0.9,p = 10^-10)
```

Arguments

| | |
|-------|---|
| DE | A matrix with one column of zscore. |
| nperm | Number of random permutations (default: 100). |
| r | Restart the probability of the random-walk algorithm (default: 0.9). |
| p | For each node, if the difference in centrality score between iterations changes less than this value, the algorithm considers the calculation complete (default: 10^-10). |

Value

A dataframe with seven columns those are drugbankid, centralscore, p.value,fdr,number of targets, drug targets,drugname.

Examples

```
# Obtain the example data
GEP<-Gettest("GEP")
label<-Gettest("label")
# Run the function
DEscore<-CalDEscore(GEP,label)
# Run the function
drug_centrality<-DrugReposition(DE=DEscore,nperm = 100,r = 0.9,p = 10^-10)
```

*DrugSimscore**DrugSimscore*

Description

The function "DrugSimscore" is used in calculating the drug functional similarity score.

Usage

```
DrugSimscore(DE,nperm = 0)
```

Arguments

- DE A matrix with one column of zscore.
 nperm Number of random permutations (default: 0).

Value

A dataframe with four columns those are drug1, drug2, drug1 name, drug2 name, functional similarity score and FDR.

Examples

```
# Obtain the example data
GEP<-Gettest("GEP")
label<-Gettest("label")
# Run the function
DEscore<-CalDEscore(GEP,label)
# Run the function
drug_drug<-DrugSimscore(DE=DEscore,nperm = 0)
```

Gettest

*Gettest***Description**

Get the example data

Usage

```
Gettest(exampleData)
```

Arguments

- exampleData A character,should be one of "Jaccard", "commongenes", "GO_MF", "Drugs", "Drugbankid_CID", "drug

Value

data

myenv

*An environment variable which includes some example data***Description**

An environment variable which includes some example data. Jaccard:A matrix of Jaccard score between drugs and GOMF. commongenes:A matrix consisting of genes shared by drug targets and GOMF. GO_MF:GO terms of molecular functions. Drugs:Drugs and corresponding targets. GEP:An example gene expression profile. label:A vector representing the label of the sample of GEP, where "1" is the disease sample and "0" is the normal sample. Drugbankid_CID:A dataframe including three columns which are drugbankid, ChembleID, and drugname.

Usage

```
myenv
```

Format

An environment variable

```
plotDruglink
```

```
plotDruglink
```

Description

The function "plotDruglink" is used to plot a bipartite network of drugs and shared molecular functions.

Usage

```
plotDruglink(drug1,drug2,i = 5,color_MF = "#43AAEF",color_drug = "#F7525B",
layout_type = "circle")
```

Arguments

| | |
|-------------|--|
| drug1 | The drugbank ID of drug1. |
| drug2 | The drugbank ID of drug2. |
| i | Specifies the number of outputs molecular functions, which is 5 by default. |
| color_MF | Defines the color of MF nodes in the network. |
| color_drug | Defines the color of drug nodes in the network. |
| layout_type | layout_type used to set the appropriate arrangement, there is an option to choose from "circle","dh",and "sugiyama". |

Value

A bipartite network of drugs and shared molecular functions.

Examples

```
# Set drug1
drug1<- "DB02721"
# Set drug2
drug2<- "DB01213"
# Run the function
library(igraph)
plotDruglink(drug1,drug2,i = 5)
```

`plotDrugstructure` *plotDrugstructure*

Description

The function "plotDrugstructure" can plot the chemical structure of a drug.

Usage

```
plotDrugstructure(drugid = "")
```

Arguments

`drugid` A drugbank ID.

Value

A chemical structure of specific drug

Examples

```
# Load depend package
library(ChemmineR)
library(rvest)
# Obtain molecular formula and visualize it.
plotDrugstructure(drugid ="DB00780")
```

`plotTargetheatmap` *plotTargetheatmap*

Description

The function "plotTargetheatmap" is used to plot a heat map of drug targets expression.

Usage

```
plotTargetheatmap(drugid,ExpData,label,significance=FALSE,
cluster.rows=FALSE,cluster.cols=FALSE,bk=c(-2.4,2.3),show.rownames=TRUE,
show.colnames=FALSE,ann_colors=c("#FFAA2C","#2CBADA"),col=c("#2A95FF","#FF1C1C"))
```

Arguments

| | |
|---------------------------|---|
| <code>drugid</code> | The drugbank ID of a drug. |
| <code>ExpData</code> | A gene expression profile of interest (rows are genes, columns are samples). |
| <code>label</code> | A character vector consists of "0" and "1" which represent sample class in the gene expression profile. "0" means normal sample and "1" means disease sample. |
| <code>significance</code> | This parameter controls whether the p-value of differential expression is displayed. |

| | |
|---------------|--|
| cluster.rows | Logical value that represents whether row clustering is used. |
| cluster.cols | Logical value that represents whether col clustering is used. |
| bk | This parameter adjusts the range of values displayed by the color bar. |
| show.rownames | This parameter controls whether row names are displayed. |
| show.colnames | This parameter controls whether column names are displayed. |
| ann_colors | Vector of colors used to define groups. |
| col | Vector of colors used in the heatmap. |

Value

A heat map of drug targets expression.

Examples

```
# Obtain the example data
GEP<-Gettest("GEP")
label<-Gettest("label")
# Run the function
plotTargetheatmap("DB00780",GEP,label)
```

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