Package ‘DrInsight’

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Title Drug Repurposing: Integration and Systematic Investigation of Genomic High Throughput Data

Version 0.1.1

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Description This is a connectivity mapping-based drug repurposing tool that identifies drugs that can potentially reverse query disease phenotype or have similar functions with query drugs.

Depends R (>= 3.4.0), igraph (>= 1.1.2), qusage (>= 2.12.0)

License GPL-2

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**Description**

This data set gives the TCGA breast cancer tumor versus normal t-test scores of each gene.

**Usage**

`brca.tcg`a

**Format**

A matrix containing the t statistics of 17960 genes

**Source**

TCGA
**drug.ident**

**Arguments**

path.analysis.res  
The pathway analysis results. Output of function pathway.analysis().

pathway.FDR.cutoff  
The FDR threshold to select significant drug specific pathways and the default is 0.1.

drug.name  
The name of the drug user would like to analyze. The specified drug name should among those that are listed in the output table of function network.graph().

pathway.name  
The name of the pathway user would like to analyze. The specified pathway name should among those that are listed in the output table of function network.graph().

show.plot  
True or False, specifying if you want to show the pathway-CEG plot for a specific drug and pathway.

**Examples**

```r
## get the Dr. Insight drug identification results
drug.ident.res = drug.ident(query.data = example.disease, cmap.ref.profiles = example.drug.profiles, repurposing.unit = "treatment", connectivity = "negative")

## load in example pathway data
data("example.pathway")

## Performe pathway analysis (for the drugs that are identified by ident.drug())
path.analysis.res = pathway.analysis(drug.ident.res = drug.ident.res, pathway.list = example.pathway, drug.FDR.cutoff = 0.5)

path.CEG.network = CEG.network.graph(path.analysis.res = path.analysis.res, pathway.FDR.cutoff = 0.5, drug.name = "drug1", pathway.name = "pathway5", show.plot = TRUE)
```

---

**Description**

This function allows user to use the differential expression data of their interested disease phenotype or drug profile to query against CMap reference drug expression profiles. This function returns an object of three elements: a table of drug p-values reflecting the connectivity between query data and CMap drugs; a table contains all the meta information of CMap drugs; and an object of p-values of identified CEGs for each drug instance.

**Usage**

```r
drug.ident(query.data = NULL, cmap.ref.profiles = NULL, repurposing.unit = "treatment", CEG.threshold = 0.05, connectivity = "negative")
```
Arguments

query.data  User provided differential expression analysis results (e.g. t-test statistic scores) of querying data (either disease phenotype data, or drug expression data). The column names for gene symbols and statistic scores must be "geneSymbol" and "score".

cmap.ref.profiles  Dr. Insight provided CMap drug rank matrix containing all 6100 instances of CMap data set. The example of reference data can be loaded with data("example.profiles"). The real cmap data can be loaded with the function 'get.cmap.ref' (See instructions in vignette).

repurposing.unit  The parameter of either "treatment" or "drug", which indicates if user want the algorithm to test drug repurposing p value at treatment level or drug level. The default is "treatment", which treats the drug data from different cell lines separately.

CEG.threshold  The p value threshold to select the consistently differential expressed genes (CEGs).

connectivity  The type of connectivity, either "negative" or "positive". Negative connectivity is used when the query data is the differential scores from disease data, and Dr. Insight will repurpose drugs that can potentially reverse the query disease phenotype. Positive connectivity is used when the query data is from a drug profile, and Dr. Insight will return the drugs that are similar to the query drug.

Examples

```r
## load in the example query disease data
data("example.disease")
data("example.drug.profiles")

## get the Dr. Insight drug identification results
drug.ident.res = drug.ident(query.data = example.disease, cmap.ref.profiles = example.drug.profiles, repurposing.unit = "treatment", connectivity = "negative")

drug.pvals = drug.ident.res$drug.pvals
```

drug.info  *The matrix containing the drug instances information of CMap drug data*

Description

This matrix gives the drug instance information of CMap drug data, including drug names, instance numbers, cell line, dosage, etc.

Usage

drug.info
example.disease

Format
A matrix of 6100 rows and 7 columns.

Source
CMap

| example.disease | An example gene differential expression analysis test statistic score data |

Description
This data gives an example of gene differential expression analysis test statistic score as the input for DrInsight analysis.

Usage
example.disease

Format
A matrix containing the simulated scores of 1000 genes

Source
simulation

| example.drug.profiles | An example cmap drug reference profiles |

Description
This data gives an simulated example of cmap drug perturbed reference profiles.

Usage
example.drug.profiles

Format
An object containing the simulated reference drug rank matrix and drug information table

Source
simulation
example.pathway  example.pathway

Description
This data set gives the geneset list of a simulated pathways.

Usage
eexample.pathway

Format
A list containing 10 pathways.

Source
simulation

get.cmap.ref  Read In and Process CMap Reference Profiles

Description
This function allows user to load in the CMap drug rank matrix user downloaded from CMap website: https://portals.broadinstitute.org/cmap/ (data matrix in the "downloads" section).

Usage
get.cmap.ref(cmap.data.path = NULL, probe.to.genes = NULL, drug.info = NULL)

Arguments
cmap.data.path  The local path and the name of the downloaded data matrix file. The data matrix file should be in txt format.

probe.to.genes  The ID converter between Affymetrix probe IDs (the IDs used in CMap data matrix) and official gene symbol. The packge comes with an embeded probe.to.genes file that can be directly used. User can use their own converter file with two columns named "ID", and "Gene.Symbol".

drug.info  The drug instance information from CMap data. The deafault drug.info file comes with DrInsight package.
make.cytoscape.network

*Drug and Pathway connection output files for Cytoscape visualization*

**Description**

This function allows user to get the two files needed for Cytoscape to visualize the drug-pathway network.

**Usage**

```r
make.cytoscape.network(path.analysis.res = path.analysis.res,
                        pathway.FDR.cutoff = 0.1)
```

**Arguments**

- `path.analysis.res`  
  The pathway analysis results. Output of pathway.analysis().

- `pathway.FDR.cutoff`  
  The FDR threshold to select significant drug specific pathways and the default is 0.1.

**Examples**

```r
## get the Dr. Insight drug identification results
drug.ident.res = drug.ident(query.data = example.disease, cmap.ref.profiles = example.drug.profiles,
                           repurposing.unit = "treatment", connectivity = "negative")

## load in example pathway data
data("example.pathway")

## Performe pathway analysis (for the drugs that are identified by ident.drug())
path.analysis.res = pathway.analysis(drug.ident.res = drug.ident.res,
                                    path.widget = example.pathway, drug.FDR.cutoff = 0.5)

## get the pathway analysis output that can be loaded into Cytoscape for visualization
network.cytoscape = make.cytoscape.network(path.analysis.res = path.analysis.res,
                                            pathway.FDR.cutoff = 0.5)
```
network.n.graph

Plot Drug and Pathway Interaction Network

Description

This function allows user to plot the drug-pathway interaction network and returns the drug-pathway interaction contingency table.

Usage

network.n.graph(path.analysis.res = path.analysis.res,
    pathway.FDR.cutoff = 0.1, pathway.label = FALSE, drug.label.size = 1.2,
    path.label.size = 0.8, return.adj.table = TRUE)

Arguments

- `pathway.FDR.cutoff`: The FDR threshold to select significant drug specific pathways and the default is 0.1.
- `pathway.label`: True or False specifying if show pathway labels in the graph.
- `drug.label.size`: The number indicating the size of drug labels in the graph.
- `path.label.size`: The number indicating the size of pathway labels in the graph.
- `return.adj.table`: True or False specifying if return the resulted drug-pathway contingency table. In the returned table, 1's represent that the pathway is up-regulated by the drug; -1's represent down-regulation and 0's no-regulation.

Examples

```r
## get the Dr. Insight drug identification results
drug.ident.res = drug.ident(query.data = example.disease, cmap.ref.profiles = example.drug.profiles,
    repurposing.unit = "treatment", connectivity = "negative")

## load in example pathway data
data("example.pathway")

## Performe pathway analysis (for the drugs that are identified by ident.drug())
path.analysis.res = pathway.analysis(drug.ident.res = drug.ident.res,
    pathway.list = example.pathway, drug.FDR.cutoff = 0.5)

drug.pathway.network = network.graph(path.analysis.res, pathway.FDR.cutoff = 0.5,
    return.adj.table = TRUE, pathway.label = TRUE)
```
**Description**

This function allows users to run pathway analysis on the identified significant drugs, therefore to detect drug mechanism related pathways, and the CEGs in the pathways.

**Usage**

```r
pathway.analysis(drug.ident.res = drug.ident.res, pathway.list = NULL,
                  pathway.list.path = NULL, drug.FDR.cutoff = 0.1, CEG.threshold = 0.05)
```

**Arguments**

- `drug.ident.res` Dr. Insight drug identification analysis results. Output of function `drug.ident()`.
- `pathway.list` The pathways used to analyze drug mechanism. Should be a list of pathways where the names of lists are pathway names. Dr.Insight provides NIH pathway interaction database (PID) pathways, which can be called by `data("pathway.PID")`. If you want to analyze with other pathways (from GSEA MsigDB), please use the parameter: `pathway.list.path`.
- `pathway.list.path` The local path and file name of the pathways downloaded from GSEA MsigDB http://software.broadinstitute.org/gsea/msigdb/collections.jsp, if `pathway.list` is NULL. Should be in full directory and file name (See instructions in vignette).
- `drug.FDR.cutoff` The FDR threshold to select significant repurposed drugs by Dr. Insight for pathway analysis.
- `CEG.threshold` The p value threshold for selecting significant consistently differential expressed genes (CEGs).

**Examples**

```r
## get the Dr. Insight drug identification results
drug.ident.res = drug.ident(query.data = example.disease, cmap.ref.profiles = example.drug.profiles,
                   repurposing.unit = "treatment", connectivity = "negative")

## load in example pathway data
data("example.pathway")

## Performe pathway analysis (for the drugs that are identified by ident.drug())
path_analysis.res = pathway.analysis(drug.ident.res = drug.ident.res,
                                      pathway.list = example.pathway, drug.FDR.cutoff = 0.5)
```
### pathway.Npid

**Description**

This data set gives the geneset list of PID pathways

**Usage**

pathway.Npid

**Format**

A large list containing 222 pathways.

**Source**

PID genesets from Pahtway Commons

---

### probe.to.genes

**Description**

This matrix gives the resource to convert the Affymetrix probe IDs in CMap data into gene symbols.

**Usage**

probe.to.genes

**Format**

A matrix of two columns and 12994 rows

**Source**

Affymetrix
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