

Package ‘immcp’

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Title Poly-Pharmacology Toolkit for Traditional Chinese Medicine Research

Version 1.0.3

Description Toolkit for Poly-pharmacology Research of Traditional Chinese Medicine. Based on the biological descriptors and drug-disease interaction networks, it can analyze the potential poly-pharmacological mechanisms of Traditional Chinese Medicine and be used for drug-repositioning in Traditional Chinese Medicine.

License GPL (>= 3)

URL <https://github.com/YuanlongHu/immcp>

BugReports <https://github.com/YuanlongHu/immcp/issues>

Depends igraph, R (>= 4.0.0)

Imports clusterProfiler, DOSE, dplyr, methods, magrittr, Matrix, openxlsx, org.Hs.eg.db, pbapply, proxyC, purrr, rlang (>= 1.0.2), stats, utils, visNetwork (>= 0.3.1), arules, ggplot2, ggheatmap, factoextra

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BasicData-class	<i>Class BasicData This class represents the basic input data.</i>
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Description

Class BasicData This class represents the basic input data.

Slots

- drugnet A directed graph
- vertices Vertices of drug graph.
- diseasenet Disease network.
- biomarker Disease-related gene.

Author(s)

Yuanlong Hu

BioDescr-class	<i>Class BioDescr This class represents the biological descriptor data.</i>
----------------	---

Description

Class BioDescr This class represents the biological descriptor data.

Slots

drug_geneset from drug to geneset.
 geneset_gene from geneset to gene for each drug.
 anno Geneset ID and description.

Author(s)

Yuanlong Hu

CreateBasicData	<i>CreateBasicData</i>
-----------------	------------------------

Description

Create BasicData Object

Usage

```
CreateBasicData(..., diseasenet = NULL, biomarker = NULL)
```

Arguments

...	Drug graph from PrepareData.
diseasenet	A graph of Disease-related gene from PrepareData.
biomarker	Character vector, the vector of Disease-related gene.

Value

A BasicData object.

Author(s)

Yuanlong Hu

Examples

```
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseasenet = disease)
```

CreateDisDrugNet

CreateDisDrugNet

Description

Create Disease-Drug Network

Usage

```
CreateDisDrugNet(BasicData, drug, disease)
```

Arguments

BasicData	BasicData object.
drug	Character vector, the drug.
disease	Character vector, the disease.

Value

A igraph object.

Author(s)

Yuanlong Hu

Examples

```
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseasenet = disease)
DisDrugNet <- CreateDisDrugNet(BasicData, drug = "Drug1", disease = "disease")
```

diff_network_char	<i>diff_network_char</i>
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Description

Calculate the difference of network characters in two network

Usage

diff_network_char(graph1, graph2, output_all = FALSE)

Arguments

- graph1 A igraph object.
- graph2 A igraph object.
- output_all FALSE

Value

A number vector.

Author(s)

Yuanlong Hu

drugdemo	<i>Datasets Demo dataset</i>
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Description

Datasets Demo dataset

enrich_f	<i>enrich_f</i>
----------	-----------------

Description

Enrich Analysis

Usage

```
enrich_f(  
  target_character,  
  geneset = c("kegg", "mkegg", "go", "wp"),  
  arguments = list(minGSSize = 5, maxGSSize = 500, pvalue = 0.05, qvalue = 0.1),  
  out_dataframe = TRUE,  
  to_ENTREZID = TRUE  
)
```

Arguments

target_character	Charactor vector of gene.
geneset	Charactor vector, one of "kegg"(KEGG), "mkegg"(KEGG Module), "go"(GO-BP), and "wp"(WikiPathways); a data frame and list.
arguments	A list of the arguments of clusterProfiler, including minGSSize, maxGSSize, pvalue, and qvalue.
out_dataframe	Logical, whether to output data frame,defaults to FALSE.
to_ENTREZID	Logical, whether to translate to ENTREZID from SYMBOL, defaults to TRUE.

Value

data frame

Author(s)

Yuanlong Hu

exportCytoscape	<i>Export an xlsx file to Cytoscape</i>
-----------------	---

Description

Export an xlsx file to Cytoscape.

Usage

```
exportCytoscape(graph, file)
```

Arguments

graph	igraph object.
file	file

Value

A workbook object

Author(s)

Yuanlong Hu

extr_biodescr	<i>Extract Biological descriptor</i>
---------------	--------------------------------------

Description

Extract Biological descriptor

Usage

```
extr_biodescr(  
  BasicData,  
  geneset = c("kegg", "mkegg", "go", "wp"),  
  arguments = list(minGSSize = 5, maxGSSize = 500, pvalue = 0.05, qvalue = 0.1),  
  ref_type = "drug",  
  ref = NULL,  
  to_ENTREZID = TRUE  
)
```

Arguments

BasicData	BasicData object.
geneset	Character vector, one of "kegg"(KEGG), "mkegg"(KEGG Module), "go"(GO-BP), and "wp"(WikiPathways); a data frame and list.
arguments	A list of the arguments of clusterProfiler, including minGSSize, maxGSSize, pvalue, and qvalue.
ref_type	Character vector, one of "drug", "herb", "compound" or "target", defaults to "drug".
ref	Character vector, reference drug, herb, compound or target, defaults to NULL.
to_ENTREZID	Logical, whether to translate to ENTREZID from SYMBOL, defaults to TRUE.

Value

A BioDescr object.

Author(s)

Yuanlong Hu

extr_biodescr, BasicData-method
Extract Biological descriptor

Description

Extract Biological descriptor

Usage

```
## S4 method for signature 'BasicData'
extr_biodescr(
  BasicData,
  geneset = c("kegg", "mkegg", "go", "wp"),
  arguments = list(minGSSize = 5, maxGSSize = 500, pvalue = 0.05, qvalue = 0.1),
  ref_type = "drug",
  ref = NULL,
  to_ENTREZID = TRUE
)
```

Arguments

BasicData	BasicData object.
geneset	Character vector, one of "kegg"(KEGG), "mkegg"(KEGG Module), "go"(GO-BP), and "wp"(WikiPathways); a data frame and list.

arguments	A list of the arguments of clusterProfiler, including minGSSize, maxGSSize, pvalue, and qvalue.
ref_type	Character vector, one of "drug", "herb", "compound" or "target", defaults to "drug".
ref	Character vector, reference drug, herb, compound or target, defaults to NULL.
to_ENTREZID	Logical, whether to translate to ENTREZID from SYMBOL, defaults to TRUE.

Value

A BioDescr object.

Examples

```
## Not run:
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseaseset = disease)
biodescr <- extr_biodescr(BasicData, geneset= "kegg")

## End(Not run)
```

HerbResult-class	<i>Class HerbResult This class represents the biological descriptor data.</i>
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Description

Class HerbResult This class represents the biological descriptor data.

Slots

Drug_Herb Data frame, Drug-herb relationship.

Herb_Herb Herb-herb association Rule Graph, it is a directed graph.

Author(s)

Yuanlong Hu

natural_connectivity	<i>natural_connectivity</i>
----------------------	-----------------------------

Description

Calculate the natural connectivity

Usage

natural_connectivity(graph)

Arguments

graph A igraph object.

Value

A numeric vector.

Author(s)

Yuanlong Hu

network_char	<i>network_char</i>
--------------	---------------------

Description

Calculate the network characters

Usage

network_char(graph, total_network = FALSE)

Arguments

graph The graph.
total_network Calculate for total network or each nodes.

Value

A number vector or data frame.

Author(s)

Yuanlong Hu

network_node_ks	<i>network_node_ks</i>
-----------------	------------------------

Description

Kolmogorov-Smirnov tests for node characters between networks

Usage

```
network_node_ks(graph1, graph2, replicate = 1000)
```

Arguments

graph1	A igraph object.
graph2	A igraph object.
replicate	Number vector, the number of conduct bootstrapping sampling replications.

Value

A data frame

Author(s)

Yuanlong Hu

plot_BioDescr	<i>Plot Biological descriptor</i>
---------------	-----------------------------------

Description

Plot Biological descriptor

Usage

```
plot_BioDescr(  
  BioDescr,  
  type = "heatmap",  
  cluster_k = 2,  
  colors = c("#2E9FDF", "#E7B800")  
)
```

Arguments

BioDescr	BioDescr object.
type	one of "heatmap" and "clusterplot".
cluster_k	Number vector, number of cluster.
colors	vector of colors.

Value

Returns NULL, invisibly.

plot_graph

Plot Disease-Drug Network

Description

Plot Disease-Drug Network

Usage

```
plot_graph(  
  graph,  
  drug,  
  disease,  
  Isolated = TRUE,  
  vis = "visNetwork",  
  color = c(drug = "#cca4e3", herb = "#ff461f", compound = "#ffc773", target =  
    "#70f3ff"),  
  width = 1,  
  size = 20,  
  ...  
)  
  
## S4 method for signature 'BasicData'  
plot_graph(  
  graph,  
  drug,  
  disease,  
  Isolated = TRUE,  
  vis = "visNetwork",  
  color = c(drug = "#cca4e3", herb = "#ff461f", compound = "#ffc773", target =  
    "#70f3ff"),  
  width = 1,  
  size = 20,  
  ...  
)  
  
## S4 method for signature 'igraph'  
plot_graph(  
  graph,  
  drug,  
  disease,  
  Isolated = TRUE,  
  vis = "visNetwork",
```

```
    color = c(drug = "#cca4e3", herb = "#ff461f", compound = "#ffc773", target =
      "#70f3ff"),
    width = 1,
    size = 20,
    ...
  )

## S4 method for signature 'HerbResult'
plot_graph(
  graph,
  drug,
  disease,
  Isolated = TRUE,
  vis = "visNetwork",
  color = c(drug = "#cca4e3", herb = "#ff461f", compound = "#ffc773", target =
    "#70f3ff"),
  width = 1,
  size = 20,
  ...
)
```

Arguments

graph	graph.
drug	drug.
disease	disease.
Isolated	Whether to delect Isolated nodes.
vis	one of "igraph", "visNetwork" and "shiny".
color	Nodes Color
width	Edges width
size	Nodes size
...	Arguments

Value

Returns NULL, invisibly.

Author(s)

Yuanlong Hu

PrepareData

PrepareData

Description

Prepare input format.

Usage

```
PrepareData(..., from, to, diseaseID, format = "single", sep)
```

Arguments

<code>...</code>	data frame, containing interaction information.
<code>from</code>	A character vector, containing "drug", "herb", "compound", or "target".
<code>to</code>	A character vector, containing "drug", "herb", "compound", or "target".
<code>diseaseID</code>	Character vector, diseaseID
<code>format</code>	one of "single" or "basket".
<code>sep</code>	Separator.

Value

A igraph object.

Author(s)

Yuanlong Hu

Examples

```
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
```

`read_gmt`*write_gmt*

Description

parse gmt file to a data.frame

Usage

```
read_gmt(gmtfile, out_dataframe = TRUE)
```

Arguments

`gmtfile` A GMT file name or URL containing gene sets.
`out_dataframe` TRUE or FALSE

Value

data.frame, list

Author(s)

Yuanlong Hu

`score_network`*score_network*

Description

Calculating differences in disease network characteristics before and after removal of drug targets

Usage

```
score_network(BasicData, n = 1000)
```

Arguments

`BasicData` A BasicData object.
`n` Number vector, the number of times random permutation sampling, default to 1000.

Value

A list.

Author(s)

Yuanlong Hu

Examples

```
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseasenet = disease)
res <- score_network(BasicData, n = 100)
```

score_rule

*Mining herb-herb associations with Apriori***Description**

Mine herb-herb association rules of prescription using the Apriori algorithm.

Usage

```
score_rule(BasicData, drug = NULL, support = 0.1, confidence = 0.8)
```

Arguments

BasicData	BasicData object.
drug	Character vector of drug names to analyze, default to NULL.
support	A numeric value for the minimal support of an item set, default to 0.1.
confidence	A numeric value for the minimal confidence of an item set, default to 0.8.

Value

A HerbResult object.

Author(s)

Yuanlong Hu

Examples

```
## Not run:
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseasenet = disease)
```



```
res <- score_rule(BasicData, support = 0.1, confidence = 0.8)

## End(Not run)
```

score_sim	<i>Calculating similarity between drug and disease</i>
-----------	--

Description

Calculating drug-disease similarity based on biological descriptors

Usage

```
score_sim(BioDescr, method = "jaccard", n = 1000)
```

Arguments

BioDescr	BioDescr object.
method	method to compute similarity, default "jaccard". See <code>proxyC::simil</code> .
n	number.

Value

A list.

Author(s)

Yuanlong Hu

Examples

```
## Not run:
data(drugdemo)
drug_herb <- PrepareData(drugdemo$drug_herb, from = "drug", to="herb")
herb_compound <- PrepareData(drugdemo$herb_compound, from = "herb", to="compound")
compound_target <- PrepareData(drugdemo$compound_target, from = "compound", to="target")
disease <- PrepareData(drugdemo$disease, diseaseID = "disease", from = "target", to="target")
BasicData <- CreateBasicData(drug_herb, herb_compound, compound_target, diseasenet = disease)
biodescr <- extr_biodescr(BasicData, geneset= "kegg")
res <- score_sim(biodescr, method="jaccard", n=1000)

## End(Not run)
```

<code>to_biodescr</code>	<i>to_biodescr</i>
--------------------------	--------------------

Description

Convert BioDescr object to a list of adjacency matrix

Usage

```
to_biodescr(BioDescr)
```

Arguments

BioDescr A BioDescr object.

Value

A list.

Author(s)

Yuanlong Hu

Examples

```
## Not run:
  to_biodescr(BioDescr)

## End(Not run)
```

<code>to_df</code>	<i>to_df</i>
--------------------	--------------

Description

Convert list to data.frame

Usage

```
to_df(list)
```

Arguments

list A list containing gene sets.

Value

A data frame.

Author(s)

Yuanlong Hu

Examples

```
## Not run:  
to_df(list)  
  
## End(Not run)
```

to_list

to_list

Description

Create a new list from a data.frame of drug target and disease biomarker as input

Usage

```
to_list(dataframe, input = "single", sep = ", ")
```

Arguments

dataframe	a data frame of 2 column with term/drug and gene
input	one of the single or basket
sep	When 'input' is 'basket'.

Value

list

Author(s)

Yuanlong Hu

Examples

```
## Not run:  
to_list(dataframe)  
  
## End(Not run)
```

`write_gmt`*write_gmt*

Description

prints data frame to a gmt file

Usage

```
write_gmt(geneset, gmt_file)
```

Arguments

<code>geneset</code>	A data.frame of 2 column with term/drug and gene.
<code>gmt_file</code>	A character of gmt file name.

Value

gmt file

Author(s)

Yuanlong Hu

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