Package 'FunJaccR'

March 14, 2025

Title Interpretation of Gene Lists Using Clustering of Functionally

Type Package

Enriched Terms

Version 0.1.0
Description Given a gene list, funtionally enriched terms are determined using gProfiler. MCL is used to cluster these terms based on the Jaccard similarity of the lists of genes associated with them. This can be visualised in Cytoscape using the RCy3 package.
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```
create_cytoscape_network
```

Create a FunJacc network in Cytoscape

Description

Draw the FunJacc results as a network in an open instance of Cytoscape running on the same machine

Usage

```
create_cytoscape_network(
  funjacc_results,
  title = "my first network",
  collection = "Funjacc Networks"
)
```

Arguments

```
funjacc_results
Funjacc results
title Network title
collection Network collection
```

Value

Sets up a network in your Cytoscape application

Examples

```
create_cytoscape_network(funjacc_res, title="Test network")
```

funjacc

Run FunJacc

Description

Determine enriched functional term clusters with FunJacc

Usage

```
funjacc(
  gene_list,
  p_cut = 0.01,
  jaccard_cut = 0.5,
  data_types = c("GO:BP"),
  inflation = 2,
  species = "hsapiens",
  default_node_label_size = 10,
  cluster_label_size = 20
)
```

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Arguments

gene_list List of genes in which to look for enriched terms

p_cut P-value cutoff for gProfiler results

jaccard_cut Jaccard index cutoff for identifying high scoring links between terms

data_types List of functional terms to include e.g.c('GO:BP', 'GO:MF', 'GO:CC', 'KEGG',

'REAC', 'TF', 'MIRNA', 'CORUM', 'HP', 'HPA', 'WP'). Supplying 'all' will

include all these terms

inflation Inflation parameter for MCL clustering ~0.5-2

species Organism to use for gProfiler to interpret gene list e.g. 'hsapiens', 'mmusculus'

default_node_label_size

Node size to use for Cytoscape plotting

cluster_label_size

Node size for cluster labels in Cytoscape plotting

Value

list of results elements 'annotation' = clusters and their annotation, 'network' = network of clusters

Examples

```
funjacc(gene_list, data_types='all', species='hsapiens', inflation=2)
```

funjacc_res I	Data	file	of	output	from	run_	funjacc	function	for	testing	cre-	
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ate_cytoscape_network function

Description

Data file of output from run_funjacc function for testing create_cytoscape_network function

Usage

funjacc_res

Format

An object of class list of length 2.

Author(s)

Adam Reid <ajr236@cam.ac.uk>

run_gprofiler

gene_list

Data file of gene lists for testing

Description

Data file of gene lists for testing

Usage

gene_list

Format

An object of class character of length 79.

Author(s)

Adam Reid <ajr236@cam.ac.uk>

 $run_gprofiler$

 $Run\ gProfiler$

Description

Get gProfiler results for further processing

Usage

```
run_gprofiler(gene_list, organism)
```

Arguments

gene_list List of genes in which to look for enriched terms

organism Name of species relating to gene list

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