Package ‘KEGGlincs’

Type Package

Title Visualize all edges within a KEGG pathway and overlay LINCS data [option]

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Author Shana White

Maintainer Shana White <vandersm@mail.uc.edu>

Description See what is going on ‘under the hood’ of KEGG pathways by explicitly re-creating the pathway maps from information obtained from KGML files.

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LazyData true

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Depends R (>= 3.3), KOdata, hgu133a.db, org.Hs.eg.db (>= 3.3.0)

SystemRequirements Cytoscape (>= 3.3.0), Java (>= 8)

Suggests BiocInstaller (>= 1.20.3), knitr, rmarkdown, graph

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R topics documented:

add_edge_data .......................................................... 2
cyto_vis ................................................................. 3
data_mapping_info ...................................................... 3
expand_KEGG_edges ................................................... 4
expand_KEGG_mapping .................................................. 5
generate_mappings ....................................................... 6
get_fisher_info .......................................................... 6
get_graph_object ........................................................ 7
get_KGML ................................................................. 8
add_edge_data

Annotate KEGG edge mappings with user data

Description

Add data column[s] to object created from function expand_KEGG_edges

Usage

```r
add_edge_data(expanded_edges, KEGG_mappings, user_data, data_column_no = 3,
               only_mapped = FALSE)
```

Arguments

- `expanded_edges` The data frame object generated via the function expand_KEGG_edges
- `KEGG_mappings` The data.frame object generated by the function expand_KEGG_mappings
- `user_data` A data frame where in which the first two columns contain gene symbols representing an edge and any/all other column[s] contain corresponding edge data.
- `data_column_no` The column index for desired user data to be added
- `only_mapped` A logical indicator; if set to FALSE will return 'de-novo' edges that 'exist' in data but are not documented in KEGG

Value

A data frame object with detailed KEGG edge mappings annotated with user data

Examples

```r
p53_KGML <- get_KGML('hsa04115')
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML)
p53_edges <- expand_KEGG_edges(p53_KGML, p53_KEGG_mappings)
p53_HA1E_data <- overlap_info(p53_KGML, p53_KEGG_mappings, 'HA1E',
                              data_type = '100_bing', only_mapped = FALSE)

p53_edges_HA1E_ALL <- add_edge_data(p53_edges, p53_KEGG_mappings,
                                     p53_HA1E_data, c(3, 10, 12))
p53_edges_HA1E_MAPPED <- add_edge_data(p53_edges, p53_KEGG_mappings,
                                        p53_HA1E_data, c(3, 10, 12),
                                        only_mapped = TRUE)
```
cyto_vis  

**Description**  
View the KEGG pathway in Cytoscape. With either the ’expanded edges’ or ‘stacked nodes’ layout, users can visualize and interact with the graphs [strictly] as they are documented in the most recent KGML available from KEGG. This function is a modified version of the function send2cy(), which is part of the cyREST utility functions.

**Usage**  
cyto_vis(graph_object, title = "Cytoscape Graph Window", edge_width_attribute = "summary_score", port.number = 1234)

**Arguments**  
- **graph_object**: An igraph object such as the one generated by the function `get_graph_object`
- **title**: An optional title for the graph when it is in Cytoscape
- **edge_width_attribute**: The attribute that will be used for edge width; if data is not added or the attribute is not part of the graphing information, the edge width will default to 1.
- **port.number**: The port address for Cytoscape

**Value**  
A dynamic map in Cytoscape automatically formatted for convenient viewing.

**Examples**
```r
p53_KGML <- get_KGML("hsa04115")
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML, FALSE)
nodes <- node_mapping_info(p53_KEGG_mappings)
p53_edges <- expand_KEGG_edges(p53_KGML, p53_KEGG_mappings)
edges <- edge_mapping_info(p53_edges)
p53_graph_object <- get_graph_object(nodes, edges)
```

edge_mapping_info  

**Description**  
Modify the mapping information for desired look when graphed in Cytoscape

**Usage**  
edge_mapping_info(expanded_edges, data_added = FALSE, significance_markup = FALSE)
expand_KEGG_edges

Arguments

expanded_edges  The data frame object generated via the function expand_KEGG_edges() OR has been modified by the function add_edge_data()

data_added  A logical indicator; must be set to TRUE if user data has been added (i.e. edges modified by function add_edge_data())

significance_markup  A logical indicator; if set to TRUE will color edges based on direction and significance of correlation (as determined by user-data-analysis)

Value

A data.frame object for edges that will be passed on to the function get_graph_object

Examples

p53_KGML <- get_KGML("hsa04115")
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML)

#Default; no data added to edges:
p53_edges <- expand_KEGG_edges(p53_KGML, p53_KEGG_mappings)
p53_edge_mapping_info <- edge_mapping_info(p53_edges)

#If data is added to edges as additional attribute[s]:
p53_HA1E_data <- overlap_info(p53_KGML, p53_KEGG_mappings, "HA1E", data_type = "100_bing")
p53_edges_HA1E_data_MAPPED <- add_edge_data(p53_edges, p53_KEGG_mappings, p53_HA1E_data, c(3, 10,12), only_mapped = TRUE)
p53_edge_mapping_HA1E <- edge_mapping_info(p53_edges_HA1E_data_MAPPED, data_added = TRUE)

expand_KEGG_edges  Get detailed KEGG mapping information for each relation [edge] documented in KEGG

Description

Extract relationship information from KGML object and re-map based on normalized node information

Usage

expand_KEGG_edges(KGML_file, KEGG_mappings)

Arguments

KGML_file  An object of formal class KEGGPathway
KEGG_mappings  The data.frame object generated by the function expand_KEGG_mappings
expand\_KEGG\_mappings

Value

A dataframe object with unique entry information for all edges documented in the KEGG pathway. Note that each row has a unique combination of values for (entry1, entry2, entry1symbol, entry2symbol).

Examples

```r
p53\_KGML \textarrow{<-} \text{get\_KGML("hsa04115")}
p53\_KEGG\_mappings \textarrow{<-} \text{expand\_KEGG\_mappings(p53\_KGML, FALSE)}
p53\_edges \textarrow{<-} \text{expand\_KEGG\_edges(p53\_KGML, p53\_KEGG\_mappings)}
```

---

expand\_KEGG\_mappings  Get detailed KEGG mapping information for each map entity

Description

Extract mapping information from KGML object and normalize mappings based on multi-valued name attribute

Usage

```r
expand\_KEGG\_mappings(KGML\_file, convert\_KEGG\_IDs = TRUE)
```

Arguments

- **KGML\_file**: An object of formal class KEGGPathway
- **convert\_KEGG\_IDs**: A logical indicator; if set to FALSE will run faster however genes and compounds will remain labeled via KEGG codes (compounds) or accession numbers (genes). This option must be taken into account if data is being added. For example, the genes in ‘KO\_data’ are identified by symbols, thus it is necessary to retain the default option to convert IDs to symbols when planning to add edge data of this type.

Value

A dataframe object with unique entry information for all [node] objects documented in the KEGG pathway. Note that if multiple objects (i.e. genes or compounds) have the same entryID, this indicates that they share the same node [location] in the pathway.

Examples

```r
p53\_KGML \textarrow{<-} \text{get\_KGML("hsa04115")}
p53\_KEGG\_mappings \textarrow{<-} \text{expand\_KEGG\_mappings(p53\_KGML, FALSE)}
```
**generate_mappings**  
*The ‘boilerplate’ for this package’s desired graph style*

**Description**
Generates an object that can be converted to a JSON file and subsequently applied to the graph for the markup specified by this package and the layout mirroring KEGG. Intended for use within **cyto_vis**

**Usage**
```
generate_mappings(style_name, map_edge_width, edge_width_attribute, min_score, max_score)
```

**Arguments**
- **style_name**: An argument to name style; when used inside of **cyto_vis** no name is needed
- **map_edge_width**: A logical indicator; if FALSE no continuous mapping of edge width will be applied
- **edge_width_attribute**: The attribute that will be used for edge width; if data is not added or the attribute is not part of the graphing information, the edge width will default to 1.
- **min_score**: The minimum attribute value for the column used to map edge width
- **max_score**: The maximum attribute value for the column used to map edge width

**Value**
A list that can be converted to a JSON file to apply desired style/layout in Cytoscape

**Examples**
```
style.name = "myKEGGstyle"
mappings <- generate_mappings(style.name, FALSE)
```

---

**get_fisher_info**  
*Perform Fisher’s Exact test for edges in pathway*

**Description**
Obtain a measure for strength and significance for the relationship (i.e. an edge) based on the concordance/discordance of UP-and-DOWN regulated genes shared by two different experimental gene-knockouts Intended for use within **overlap_info**

**Usage**
```
get_fisher_info(edges, method)
```
**get_graph_object**

**Arguments**

- **edges**: The set of edges to be analyzed; Although the intended use is for LINCS data overlaps, the function should work with any typical data object as long as it has columns labeled ("UP", "DOWN", "UK1_DK2", "DK1_UK2") that contain integer values.

- **method**: The method to correct/adjust p-values for multiple testing. For available methods, type `p.adjust.methods` into command prompt and press enter.

**Value**

The input edge data.frame object with additional columns containing the results of the applied statistical test

**Examples**

```r
ex.data <- data.frame("UP" = c(70,6), "DOWN" = c(8,20),
                      "UK1_DK2" = c(4,47), "DK1_UK2" = c(3, 28))
overlaps <- get_fisher_info(ex.data, method = "BH")
```

---

**Description**

Obtain a graph object in the form of an igraph with KEGG-specific graphical information

**Usage**

```r
get_graph_object(node_mapping_info, expanded_edges, layered_nodes = FALSE)
```

**Arguments**

- **node_mapping_info**: The data.frame object generated by the function `node_mapping_info()`
- **expanded_edges**: The data.frame object generated by the function `edge_mapping_info()`
- **layered_nodes**: A logical indicator; if set to TRUE will create a graph with 'stacked' nodes that the user can manipulate when multiple nodes are mapped to one location

**Value**

A list object with the node and edge information from the graph required for mapping.

**Examples**

```r
p53_KGML <- get_KGML("hsa04115")
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML)

p53_node_mapping_info <- node_mapping_info(p53_KEGG_mappings)
p53_edge_mapping_info <- expand_KEGG_edges(p53_KGML, p53_KEGG_mappings)
```
```
#Default graph object will have 'expanded edges':
expanded_edges_graph_object <- get_graph_object(p53_node_mapping_info, 
                                              p53_edge_mapping_info)

#Graph with layered nodes:
layered_nodes_graph_object <- get_graph_object(p53_node_mapping_info, 
                                              p53_edge_mapping_info, 
                                              layered_nodes = TRUE)
```

---

**get_KGML**  
*Download and parse KGML file*

---

**Description**

Download and parse KGML file

**Usage**

`get_KGML(pathwayid, get_if_no_edges = FALSE)`

**Arguments**

- `pathwayid`: A KEGG pathway ID of the form "hsa12345" (only human pathways currently)
- `get_if_no_edges`: A logical indicator; if pathway has no edges returns null value if set to TRUE

**Value**

an object of Formal class KEGGPathway

**Examples**

```r
mtor_KGML <- get_KGML("hsa04150")

# Some pathways contain only node information; since the purpose of this
# package is to explore pathways in an edge-focused manner, the default
# options return a warning message instead of a parsed KGML file if the
# input pathway has no edges.
ribosome_KGML <- get_KGML("hsa03020")
ribosome_KGML <- get_KGML("hsa03020", get_if_no_edges = TRUE)
```

---

**KEGGlincs**  
*KEGGlincs: an R package designed to explore the edges in KEGG pathways*

---

**Description**

KEGGlincs: an R package designed to explore the edges in KEGG pathways
**Description**

Combines all other package functions for one-step pathway visualization

**Usage**

`KEGG_lincs(pathwayid, cell_line = NA, refine_by_cell_line = NA, add_L1000_edge_data = TRUE, significance_markup = TRUE, data_type = "100_full", pert_time = 96, only_mapped = TRUE, layered_nodes = FALSE, graph_title = "default", get_data = FALSE, convert_KEGG_IDs = TRUE)`

**Arguments**

- **pathwayid**: A KEGG pathway ID of the form "hsa12345" (only human pathways currently)
- **cell_line**: If left as NA will generate a pathway map without data-dependent attributes (such as edge width). To use in combination with LINCS data, choose from the set of cell lines: (A375,A549,ASC,HA1E,HCC515,HEK293T,HEKTE,HEPG2,HT29,MCF7,NCIH7SHSY5Y,SKL,SW480,VCAP)
- **refine_by_cell_line**: A logical indicator
- **add_L1000_edge_data**: A logical indicator
- **significance_markup**: A logical indicator; if set to TRUE will color edges based on direction and significance of correlation (as determined by user-data-analysis)
- **data_type**: Choose from data types: (100_full, 100_bing, 50_lm)
- **pert_time**: Choose from (6,24,48,96,120,144,168)
- **only_mapped**: A logical indicator; if set to FALSE will return ‘de-novo’ edges that ‘exist’ in data but are not documented in KEGG
- **layered_nodes**: A logical indicator; if set to TRUE will create a graph with ‘stacked’ nodes that the user can manipulate when multiple nodes are mapped to one location
- **graph_title**: An optional user-specified graph title
- **get_data**: A logical indicator; if set to true, will return the ‘expanded’ edge information for the specified pathway
- **convert_KEGG_IDs**: A logical indicator; if set to TRUE KEGG compounds will remain labeled via KEGG codes (do not need KEGGREST)

**Value**

A dynamic map in Cytoscape automatically formatted for convenient viewing and, if indicated by user, a data.frame object with detailed information for ‘expanded’ KEGG edges
Examples

## Not run:

# Default KEGG pathway with colored edges representing type of relationship:
KEGG_lincs("hsa04115", convert KEgg IDs = FALSE)

# KEGG pathway with edge width and color based on observed experimental data:
KEGG_lincs("hsa04115", "HA1E")

# Have edge information data.frame to be output to the global environment:
p53_edge_info <- KEGG_lincs("hsa04115", graph title = "p53"
                                convert KEgg IDs = FALSE, get data = TRUE)

## End(Not run)

node_mapping_info

Prepare nodes for mapping

Description

Modify the mapping information for desired look when graphed in Cytoscape

Usage

node_mapping_info(KEGG_mappings)

Arguments

KEGG_mappings The data.frame object generated by the function expand KEgg_mappings()  

Value

A data.frame object for nodes that will be passed on to the function get.graph_object

Examples

p53_KGML <- get.KGML("hsa04115")
p53.KEGG_mappings <- expand.KEGG_mappings(p53_KGML, FALSE)
p53_node_mapping_info <- node_mapping_info(p53.KEGG_mappings)
Get overlap information for pairs of gene knock-outs from LINCS data

Usage

```
overlap_info(KGML_file, KEGG_mappings, cell_type, data_type = "100_full",
pert_time = 96, only_mapped = TRUE, affy_based = FALSE,
keep_counts_only = TRUE, add_fisher_information = TRUE,
p.adjust.method = "BH")
```

Arguments

- **KGML_file**: An object of formal class KEGGPathway
- **KEGG_mappings**: The data.frame object generated by the function `expand_KEGG_mappings`
- **cell_type**: Choose from the set of cell lines: (A375,A549,ASC,HA1E,HCC515,HEK293T,HEKTE,HEPG2,HT2,SHSY5Y,SKL,SW480,VCAP)
- **data_type**: Choose from data types: (100_full, 100_bing, 50_lm)
- **pert_time**: Choose from (6,24,48,96,120,144,168)
- **only_mapped**: A logical indicator; if set to FALSE will return 'de-novo' edges that 'exist' in data but are not documented in KEGG
- **affy_based**: A logical indicator; if set to TRUE will return lists/counts based on probeID instead of gene symbol.
- **keep_counts_only**: A logical indicator; if set to FALSE will return data frame with lists [of gene symbols or probe ids] as well as counts
- **add_fisher_information**: A logical indicator; by default the relationships are analyzed for strength of correlation via Fisher's Exact Test
- **p.adjust.method**: For available methods, type `p.adjust.methods` into command prompt and press enter.

Value

A data frame where each row corresponds to information for pairs of experimental gene knock-outs from LINCS data (found in selected pathway).

Examples

```
p53_KGML <- get_KGML("hsa04115")
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML)
p53_edges <- expand_KEGG_edges(p53_KGML, p53_KEGG_mappings)

summary <- path_genes_by_cell_type(p53_KEGG_mappings)
p53_HA1E_data <- overlap_info(p53_KGML, p53_KEGG_mappings,
                          "HA1E", data_type = "100_bing",
                          only_mapped = FALSE)
```
**path_genes_by_cell_type**  
*See how many pathway gene knock-outs are available from data*

**Description**  
Check quantity of data across cell lines available from LINCS corresponding to the pathway of interest

**Usage**  
```
path_genes_by_cell_type(KEGG_mappings, pert_time = 96, get_KOs = FALSE, generate_plot = TRUE)
```

**Arguments**

- `KEGG_mappings`: The data.frame object generated by the function `expand_KEGG_mappings`
- `pert_time`: Choose from (6, 24, 48, 96, 120, 144, 168)
- `get_KOs`: Logical indicator to have data frame returned
- `generate_plot`: Logical indicator to generate histogram

**Value**  
A plot depicting percentage of pathway genes knocked-out by cell line and a data frame object listing the genes [by cell line]

**Examples**
```
p53_KGML <- get_KGML("hsa04115")
p53_KEGG_mappings <- expand_KEGG_mappings(p53_KGML)
path_genes_by_cell_type(p53_KEGG_mappings)
```

**refine_mappings**  
*Refine pathway by cell type*

**Description**  
Reduce the KEGG pathway by only including genes that are expressed within a given cell type

**Usage**  
```
refine_mappings(KEGG_mappings, cell_line)
```

**Arguments**

- `KEGG_mappings`: The data.frame object generated by the function `expand_KEGG_mappings`
- `cell_line`: Choose from the set of cell lines with baseline data; cell-lines may or may not have corresponding KO data
**toCytoscape**

**Description**
A subset of the R utility functions available from/defined by cyREST. The function mapAttributes is called from within toCytoscape which, in turn, is called from within cyto_vis.

**Usage**

toCytoscape(igraphobj)

mapAttributes(attr.names, all.attr, i)

**Arguments**

- **igraphobj** A graph object compatible for use with the package igraph
- **attr.names** Attribute names of an igraph object
- **all.attr** The attribute value if an igraph object
- **i** The index for a given igraph object

**Value**
A JSON object to be sent to Cytoscape
Index

add_edge_data, 2

cyto_vis, 3, 6

dge_mapping_info, 3
expand_KEGG_edges, 4
expand_KEGG_mappings, 5

generate_mappings, 6
get_fisher_info, 6
get_graph_object, 3, 7
get_KGML, 8

KEGG_lincs, 9
KEGGlincs, 8
KEGGlincs-package (KEGGlincs), 8

mapAttributes(toCytoscape), 13

node_mapping_info, 10

overlap_info, 6, 11

path_genes_by_cell_type, 12

refine_mappings, 12

toCytoscape, 13