NCIgraph

October 25, 2011

NCI.demo.cyList 10 raw NCI networks from Nature curated pathways and BioCarta imported

Description

These are the ten first elements of the full list of raw networks that can be downloaded using the downloadCyLists.R script.

Usage

NCI.demo.cyList

Format

A list of 10 graphNEL objects.

Author(s)

Laurent Jacob

Examples

```
data("NCIgraphVignette")
length(NCI.demo.cyList)
library(Rgraphviz)
plot(NCI.demo.cyList[[1]])
```

NCIgraph

Description

Package: NCIgraph Class NCIgraph

public static class **NCIgraph** extends graphNELObject

Class extending graphNEL fro graphs build from NCI gene networks.

Author(s)

Laurent Jacob

directedBFS Uses a breadth first search on a directed graph to identify which genes

Description

Uses a breadth first search on a directed graph to identify which genes are regulated by a particular node in the graph.

Usage

directedBFS(g, node)

Arguments

g	A graph object.
node	A node of g.

Value

A structured list containing the regulated genes and the type of interaction between node and each gene.

Author(s)

Laurent Jacob

See Also

propagateRegulation()

edgesToMerge Identifies edges that should be merged to parse a NCI network

Description

Identifies edges that should be merged to parse a NCI network.

Usage

```
edgesToMerge(g)
```

Arguments

g A graph object.

Value

A list of edges to be merged

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

getNCIPathways Loads networks from Cytoscape and parses them

Description

Loads networks from Cytoscape and parses them.

Usage

```
getNCIPathways(cyList=NULL, parseNetworks=TRUE, entrezOnly=TRUE, verbose=FALSE)
```

Arguments

cyList	a list providing the networks loaded from Cytoscape. If NULL, the function will try to build the list from Cytoscape.
verbose	If TRUE, extra information is output.
parseNetworks	
	A logical. If FALSE, the raw NCI networks are returned as graphNEL objects. If TRUE, some additional parsing is performed by the parseNCInetwork function.
entrezOnly	A logical. If TRUE, only keep nodes with an entrezID property.

Value

A list of two elements: pList, a list of graphNEL objects, and failedW a list containing the names of the networks that R failed to read from cytoscape.

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

Examples

##-----## Load NCIgraph ##-----

library(NCIgraph)

##------## Example 1: with Cytoscape ##------

Must have Cytoscape running with some networks open and Cytoscape RPC plugin started.

In this case, getNCIPathways will both read the raw networks from Cytoscape and parse

Not run:
grList <- getNCIPathways(cyList=NULL, parseNetworks=TRUE, entrezOnly=TRUE, verbose=TRUE)\$</pre>

End(Not run)

##------## Example 2: without Cytoscape ##------

Get some raw networks

data("NCIgraphVignette", package="NCIgraph")

```
## When passed a non null cyList argument (a list of networks),
## getNCIPathways will simply parse the list of networks
```

grList <- getNCIPathways(cyList=NCI.demo.cyList, parseNetworks=TRUE, entrezOnly=TRUE, ver

getSubtype.NCIgraph

Returns a list of @KEGGEdgeSubType objects describing each edge of the

Description

Returns a list of @KEGGEdgeSubType objects describing each edge of the NCI network.

is.NCIgraph

Usage

getSubtype.NCIgraph(object)

Arguments

object An NCIgraph object.

Value

A list of KEGGEdgeSubType objects.

Author(s)

Laurent Jacob

Examples

##-----## Load NCIgraph ##----library(NCIgraph) # # ------## Get some raw networks ##----data("NCIgraphVignette", package="NCIgraph") # # -----## Parse them ##----grList <- getNCIPathways(cyList=NCI.demo.cyList, parseNetworks=TRUE, entrezOnly=TRUE, ver ##------## ## Get the subtype of the second network. Some activation and some ## inhibition edges. ## ##----getSubtype.NCIgraph(grList[[2]])

```
is.NCIgraph
```

Assess whether a graph is a NCI graph

Description

Assess whether a graph is a NCI graph.

Usage

is.NCIgraph(gr)

Arguments gr

A graph object.

Value

A logical, TRUE if the graph is a NCI graph, FALSE otherwise.

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

mergeNodes

Merges a given list of nodes in a graph

Description

Merges a given list of nodes in a graph.

Usage

mergeNodes(g, mEdges, separateEntrez=TRUE, entrezOnly=TRUE)

Arguments

```
gA graph object.mEdgesA list of nodes to be merged.separateEntrezA logical. If TRUE, don't merge two nodes with entrezID.entrezOnlyA logical. If TRUE, only keep nodes with an entrezID property.
```

Value

The updated graph object

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

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parseNCInetwork Takes a NCI network and transforms it into a simpler graph only

Description

Takes a NCI network and transforms it into a simpler graph only representing inhibition/activation relationships between genes.

Usage

```
parseNCInetwork(g, propagateReg=TRUE, separateEntrez=TRUE, mergeEntrezCopies=TRU
```

Arguments

g	A graph object.	
propagateReg	A logical. If TRUE, use propagateRegulation to transform the network before parsing it.	
separateEntrez		
	A logical. If TRUE, don't merge two nodes with entrezID.	
mergeEntrezCopies		
	A logical. If TRUE, merge resulting nodes that have the same entrezID.	
entrezOnly	A logical. If TRUE, only keep nodes with an entrezID property.	

Value

The new graph object.

Author(s)

Laurent Jacob

Examples

```
## Load NCIgraph
library(NCIgraph)
## Get some raw networks
data("NCIgraphVignette", package="NCIgraph")
## Parse the first of them
parsedNetwork <- parseNCInetwork(NCI.demo.cyList[[1]],propagateReg=TRUE,separateEntrez=TF</pre>
```

```
propagateRegulation
```

Transforms the network in a way that each Biochemical Reaction node

Description

Transforms the network in a way that each Biochemical Reaction node pointing to a Complex points to what is regulated by the complex and updates the interaction types accordingly.

Usage

```
propagateRegulation(g)
```

Arguments g

A graph object.

Value

The updated graph object

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

translateNCI2GeneID

Gives the entrezID corresponding to the nodes of a graph

Description

Gives the entrezID corresponding to the nodes of a graph.

Usage

translateNCI2GeneID(g)

Arguments

g A graph object.

Value

A vector of character giving the entrez ID of the nodes of g.

translateNCI2GeneID

Author(s)

Laurent Jacob

See Also

parseNCInetwork()

Examples

```
##------
## Load NCIgraph
##------
library(NCIgraph)
## Get some raw networks
data("NCIgraphVignette", package="NCIgraph")
## Parse them
grList <- getNCIPathways(cyList=NCI.demo.cyList, parseNetworks=TRUE, entrezOnly=TRUE, ver
## Get the gene ids for the first of them
gids <- translateNCI2GeneID(grList[[1]])</pre>
```

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