Package ‘graph’

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**Description**

This generic function takes an object that inherits from the `graph` class and a node in that graph and returns a vector containing information about all other nodes that are accessible from the given node. The methods are vectorized so that `index` can be a vector.
addEdge

Usage

## S4 method for signature 'graph,character'
acc(object, index)

## S4 method for signature 'clusterGraph,character'
acc(object, index)

Arguments

- **object**: An instance of the appropriate graph class.
- **index**: A character vector specifying the nodes for which accessibility information is wanted.

Value

The methods should return a named list of integer vectors. The names of the list correspond to the names of the supplied nodes. For each element of the list the returned vector is named. The names of the vector elements correspond to the nodes that are accessible from the given node. The values in the vector indicate how many edges are between the given node and the node in the return vector.

Methods

- **object = graph**: An object of class graph.
- **object = clusterGraph**: An instance of the clusterGraph class.
- **index**: A character vector of indices corresponding to nodes in the graph.

Examples

```r
call.seed(123)
gR3 <- randomGraph(LETTERS[1:10], M<-1:2, p=.5)
acc(gR3, "A")
acc(gR3, c("B", "D"))
```

Description

A function to add an edge to a graph.

Usage

```r
addEdge(from, to, graph, weights)
```
Arguments

from    The node the edge starts at

to      The node the edge goes to.

graph   The graph that the edge is being added to.

weights A vector of weights, one for each edge.

Details

Both from and to can be vectors. They need not be the same length (if not the standard rules for replicating the shorter one are used). Edges are added to the graph between the supplied nodes.

The weights are given for each edge.

The implementation is a bit too oriented towards the graphNEL class and will likely change in the next release to accommodate more general graph classes.

If the graph is undirected then the edge is bidirectional (and only needs to be added once). For directed graphs the edge is directional.

Value

A new instance of a graph object with the same class as graph but with the indicated edges added.

Author(s)

R. Gentleman

See Also

addNode, removeEdge, removeNode

Examples

V <- LETTERS[1:4]
edL2 <- vector("list", length=4)
names(edL2) <- V
for(i in 1:4)
edL2[[i]] <- list(edges=c(2,1,2,1)[i], weights=sqrt(i))
gR2 <- graphNEL(nodes=V, edgeL=edL2, edgemode="directed")

gX <- addEdge("A", "C", gR2, 1)
gR3 <- randomEGraph(letters[10:14], .4)
gY <- addEdge("n", "l", gR3, 1)
Description

Add one or more nodes to a graph.

Usage

addNode(node, object, edges)

Arguments

node     A character vector of node names.
object   A graph
edges    A named list of edges.

Details

The supplied nodes are added to the set of nodes of the object.

If edges are provided then their must be the same number as there are nodes and the must be in the same order. The elements of the edges list are vectors. They can be character vectors of node labels for nodes in object and if so then they are added with unit weights. If the vector is numeric then it must be named (with labels corresponding to nodes in the object) and the values are taken to be the edge weights.

When the object is a distGraph then the edges must be supplied and they must contain appropriate distances for all nodes both those in object and those supplied.

Value

A new graph of the same class as object with the supplied node added to the set of nodes.

Author(s)

R. Gentleman

See Also

removeNode, removeEdge, addEdge
adj-methods

Examples

```r
V <- LETTERS[1:4]
edL1 <- vector("list", length=4)
names(edL1) <- V
for(i in 1:4)
edL1[[i]] <- list(edges=c(2,1,4,3)[i], weights=sqrt(i))
gR <- graphNEL(nodes=V, edgeL=edL1)
gX <- addNode("X", gR)

set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=.3)
g2 <- addNode("z", g1, edges=list(c("a", "h", "g")))
```

adj-methods

Methods for finding the adjacency list for selected nodes.

Description

This generic function takes an object that inherits from the graph class and a node in that graph and returns a vector containing information about all other nodes that are adjacent to the given node. This means that they are joined to the given node by an edge. The accessibility list, acc is the list of all nodes that can be reached from a specified node.

Value

The methods return vector of nodes that are adjacent to the specified node.

Methods

- `object = graph` An object that inherits from graph
- `index` An index (could be multiple) which can be either the integer offset for the node(s) or their labels.

See Also

- acc-methods

Examples

```r
set.seed(123)
gR3 <- randomGraph(LETTERS[1:4], M<-1:2, p=.5)
adj(gR3, "A")
adj(gR3, c(2,3))
```
adjacencyMatrix

Compute an Adjacency Matrix for a graphBAM object

Description
Though unwieldy for large matrices, a full adjacency matrix can be useful for debugging and export.
If the graph is “undirected” then reciprocal edges are explicit in the matrix.

Usage
adjacencyMatrix(object)

Arguments
object        A graphBAM object.

Details
Thus far only implemented for graphBAM objects.

Value
adjacencyMatrix returns an n x n matrix, where n is the number of nodes in the graph, ordered in the same manner as seen in the nodes method. All cells in the matrix are 0 except where edges are found.

Author(s)
P. Shannon

See Also
edgeMatrix

Examples
from <- c("a", "a", "a", "x", "x", "c")
to <- c("b", "c", "x", "y", "c", "a")
weight <- c(3.4, 2.6, 1.7, 5.3, 1.6, 7.9)
df <- data.frame(from, to, weight, stringsAsFactors = TRUE)
g1 <- graphBAM(df, edgemode = "directed")
adjacencyMatrix(g1)
Description

A graph representing the apoptosis pathway from KEGG, as well as a data.frame of attributes for use in plotting the graph with Rgraphviz and a list to compare the nodes with their respective LocusLink IDs.

Usage

data(apopGraph)

Details

The `apopGraph` data set contains three objects:

The first is `apopGraph`, which is an object of class `graph-NEL` and represents the hsa04210 graph from KEGG.

The second is `apopAttrs`, which is a data.frame with two columns, and a row for every node in `apopGraph`. The first column lists what color the node is represented with on the KEGG site. The second column lists the type of the node - either `genesym` or `text`. Most nodes are of type `genesym` as they represent genes, but some of the nodes in the KEGG graph were not genes and thus those nodes are of type `text`.

The third, `apopLocusLink` is a named list where the names correspond to the node names in `apopGraph`. The values of the list are the LocusLink IDs that correspond to that node in the KEGG graph.

Source


Examples

data(apopGraph)
if (require("Rgraphviz") & interactive())
  plot(apopGraph)
Class "attrData"

Description

A container class to manage generic attributes. Supports named attributes with default values with methods for vectorized access.

Objects from the Class

Objects can be created by calls of the form `new("attrData", defaults)`. The `defaults` argument should be a named list containing the initial attribute names and default values.

Slots

data: Where custom attribute data is stored

defaults: A named list of known attribute names and defualt values.

Methods

- `attrDataItem<-` signature(`self = "attrData", x = "character", attr = "character"`)
- `attrDataItem` signature(`self = "attrData", x = "character", attr = "missing"`)
- `attrDataItem` signature(`self = "attrData", x = "character", attr = "character"`)
- `attr Defaults<-` signature(`self = "attrData", attr = "character", value = "ANY"`)
- `attr Defaults<-` signature(`self = "attrData", attr = "missing", value = "list"`)
- `attr Defaults<-` signature(`self = "attrData", attr = "missing"`)
- `attr Defaults` signature(`self = "attrData", attr = "character"`)
- `initialize` signature(`.Object = "attrData"`)
- `names` return the names of the stored attributes
- `names<-` set the names of the stored attributes
- `removeAttrDataItem` signature(`self="attrData", x="character", value=NULL`): Remove the data associated with the key specified by `x`.

Author(s)

Seth Falcon

Examples

defaultProps <- list(weight=1, color="blue", friends=c("Bob", "Alice"))
adat <- new("attrData", defaults=defaultProps)

# Get all defaults
attrDefaults(adat)
## Or get only a specific attribute
attrDefaults(adat, attr="color")

## Update default weight
attrDefaults(adat, attr="weight") <- 500

## Add new attribute
attrDefaults(adat, attr="length") <- 0

## Asking for the attributes of an element you haven't customized
## returns the defaults
attrDataItem(adat, x=c("n1", "n2"), attr="length")

## You can customize values
attrDataItem(adat, x=c("n1", "n2"), attr="length") <- 5

## What keys have been customized?
names(adat)

attrDataItem-methods  Get and set attributes values for items in an attrData object

Description

The attrDataItem method provides get/set access to items stored in a attrData-class object.

Usage

attrDataItem(self, x, attr)
attrDataItem(self, x, attr) <- value

Arguments

self    A attrData-class instance
x       A character vector of item names
attr    A character vector of length 1 giving the attribute name to get/set. Note that
        the attribute name must have already been defined for the attrData object via
        attrDefaults. If missing, return a list of all attributes for the specified nodes.
value   An R object to set as the attribute value for the specified items. If the object has
        length one or does not have a length method defined, it will be assigned to all
        items in x. If the length of value is the same as x, the corresponding elements
        will be assigned. We will add an argument to indicate that the value is to be
        taken as-is for those cases where the lengths are the same coincidentally.
**attrDefaults-methods**  
*Get and set the default attributes of an attrData object*

**Description**

The `attrDefaults` method provides access to a `attrData-class` object’s default attribute list. The default attribute list of a `attrData-class` object defines what attributes can be customized for individual data elements by defining attribute names and default values.

**Usage**

```r
attrDefaults(self, attr)
attrDefaults(self, attr) <- value
```

**Arguments**

- `self`  
  A `attrData-class` instance
- `attr`  
  A character vector of length 1 giving the name of an attribute. Can be missing.
- `value`  
  An R object that will be used as the default value of the specified attribute, or a named list of attribute name/default value pairs if `attr` is missing.

**aveNumEdges**  
*Calculate the average number of edges in a graph*

**Description**

`aveNumEdges` divides the number of edges in the graph by the number of nodes to give the average number of edges.

**Usage**

```r
aveNumEdges(objgraph)
```

**Arguments**

- `objgraph`  
  the graph object

**Value**

A double representing the average number of edges will be returned.

**Author(s)**

Elizabeth Whalen
See Also

numEdges, mostEdges, numNoEdges

Examples

```r
set.seed(124)
g1 <- randomGraph(1:10, letters[7:12], p=.6)
aveNumEdges(g1)
```

biocRepos  
A graph representing the Bioconductor package repository

Description

This graph is a rendition of the Bioconductor package repository and represents the dependency graph of that repository. An edge between two package denotes a dependency on the ‘to’ package by the ‘from’ package.

Usage

data(biocRepos)

Examples

data(biocRepos)
### An example of usage will be here soon

boundary  
Returns the Boundary between a Graph and a SubGraph

Description

The boundary of a subgraph is the set of nodes in the original graph that have edges to nodes in the subgraph. The function boundary computes the boundary and returns it as a list whose length is the same length as the number of nodes in the subgraph.

Usage

```r
boundary(subgraph, graph)
```

Arguments

- **graph**: the original graph from which the boundary will be created
- **subgraph**: can either be the vector of the node labels or the subgraph itself.
The \textit{boundary} of a \textit{subgraph} is the set of nodes in the graph which have an edge that connects them to the specified subgraph but which are themselves not elements of the subgraph. For convenience users can specify the subgraph as either a graph or a vector of node labels.

\textbf{Value}

This function returns a named list of length equal to the number of nodes in \textit{subgraph}. The elements of the list correspond to the nodes in the \textit{subgraph}. The elements are lists of the nodes in \textit{graph} which share an edge with the respective node in \textit{subgraph}.

\textbf{Author(s)}

Elizabeth Whalen and R. Gentleman

\textbf{See Also}

\texttt{subGraph, graph-class}

\textbf{Examples}

```r
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=.3)
## both should be "a"
boundary(c("g", "i"), g1)
```

---

\textbf{Calculation}

\texttt{calcProb} calculates the probability of having the number of edges found in the subgraph given that it was made from \textit{origgraph}. The hypergeometric distribution is used to calculate the probability (using the pdf).

\textbf{Usage}

\texttt{calcProb(subgraph, origgraph)}

\textbf{Arguments}

\begin{itemize}
  \item \texttt{subgraph} \hspace{1em} subgraph made from the original graph
  \item \texttt{origgraph} \hspace{1em} original graph object from which the subgraph was made
\end{itemize}

\textbf{Value}

The probability of the subgraph’s number of edges is returned.
calcSumProb

Author(s)
Elizabeth Whalen

See Also
calcSumProb

Examples
#none right now

---
calcSumProb  Calculate the probability that a subgraph has an unusual number of edges.

Description
For any graph a set of nodes can be used to obtain an induced subgraph (see subGraph). An interesting question is whether that subgraph has an unusually large number of edges. This function computes the probability that a random subgraph with the same number of nodes has more edges than the number observed in the presented subgraph. The appropriate probability distribution is the hypergeometric.

Usage
calcSumProb(sg, g)

Arguments
sg  subgraph made from the original graph

Arguments
g  original graph object from which the subgraph was made

Details
The computation is based on the following argument. In the original graph there are \( n \) nodes and hence \( N = n \times (n - 1)/2 \) edges in the complete graph. If we consider these \( N \) nodes to be of two types, corresponding to those that are either in our graph, \( g \), or not in it. Then we think of the subgraph which has say \( m \) nodes and \( M = m \times (m - 1)/2 \) possible edges as representing \( M \) draws from an urn containing \( N \) balls of which some are white (those in \( g \)) and some are black. We count the number of edges in the subgraph and use a Hypergeometric distribution to ask whether our subgraph is particularly dense.

Value
The probability of having greater than or equal to the subgraph’s number of edges is returned.
Author(s)
Elizabeth Whalen

See Also
calcProb

Examples
set.seed(123)
V <- letters[14:22]
g1 <- randomEGraph(V, .2)

sg1 <- subGraph(letters[c(15,17,20,21,22)], g1)
calcSumProb(sg1, g1)

Description
This function removes all edges to or from the specified node in the graph.

Usage
clearNode(node, object)

Arguments
node a node
object a graph

Details
All edges to and from node are removed. node can be a vector.

Value
A new instance of the graph with all edges to and from the specified node(s) removed.

Author(s)
R. Gentleman

See Also
removeNode, removeEdge
Examples

```r
V <- LETTERS[1:4]
edL3 <- vector("list", length=4)
for(i in 1:4)
edL3[[i]] <- list(edges=(i%%4)+1, weights=i)
names(edL3) <- V
gR3 <- graphNEL(nodes=V, edgeL=edL3, "directed")
g4 <- clearNode("A", gR3)
```

Description

A cluster graph is a special sort of graph for clustered data. Each cluster forms a completely connected subgraph. There are no edges between clusters.

Objects from the Class

Objects can be created by calls of the form `new("clusterGraph", ...)`.

Slots

- `clusters`: Object of class "list" a list of the labels of the elements, one element of the list for each cluster.

Extends

Class "graph", directly.

Methods

- `connComp` signature(object = "clusterGraph"): find the connected components; simply the clusters in this case.
- `acc` signature(object = "clusterGraph"): find the accessible nodes from the supplied node.
- `adj` signature(object = "clusterGraph"): find the adjacent nodes to the supplied node.
- `nodes` signature(object = "clusterGraph"): return the nodes.
- `nodes<-` signature(object="clusterGraph", value="character"): replace the node names with the new labels given in value.
- `numNodes` signature(object = "clusterGraph"): return the number of nodes.
- `edgeWeights` Return a list of edge weights in a list format similar to the edges method.
- `edgeL` signature(graph = "clusterGraph"): A method for obtaining the edge list.
- `coerce` signature(from = "clusterGraph", to = "matrix"): Convert the clusterGraph to an adjacency matrix. Currently, weights are ignored. The conversion assumes no self-loops.
Description
This generic function takes an object that inherits from the `graph` class. The graph needs to have edgemode="undirected". If it has edgemode="directed", the function will return NULL.

Usage
```r
## S4 method for signature 'graph'
clusteringCoefficient(object, selfLoops=FALSE)
```

Arguments
- `object` An instance of the appropriate graph class.
- `selfLoops` Logical. If true, the calculation takes self loops into account.

Details
For a node with n adjacent nodes, if selfLoops is FALSE, the clustering coefficient is \( \frac{N}{n(n-1)} \), where N is the number of edges between these nodes. The graph may not have self loops. If selfLoops is TRUE, the clustering coefficient is \( \frac{N}{n(n+1)} \), where N is the number of edges between these nodes, including self loops.

Value
A named numeric vector with the clustering coefficients for each node. For nodes with 2 or more edges, the values are between 0 and 1. For nodes that have no edges, the function returns the value NA. For nodes that have exactly one edge, the function returns NaN.
Examples

```r
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=.3)
clusteringCoefficient(g1)
clusteringCoefficient(g1, selfLoops=TRUE)
```

Description

A collection of functions and methods to convert various forms of matrices into graph objects.

Usage

```r
aM2bpG(aM)
ftM2adjM(ft, W=NULL, V=NULL, edgemode="directed")
ftM2graphNEL(ft, W=NULL, V=NULL, edgemode="directed")
## S4 method for signature 'graphNEL,matrix'
coerce(from,to="matrix",strict=TRUE)
## S4 method for signature 'matrix,graphNEL'
coerce(from,to="graphNEL",strict=TRUE)
```

Arguments

- `ft`: An nx2 matrix containing the from/to representation of graph edges.
- `W`: An optional vector of edge weights.
- `V`: An optional vector of node names.
- `aM`: An affiliation matrix for a bipartite graph.
- `edgemode`: Character. Specifies if the resulting graph is to be directed or undirected.
- `from`: Object to coerce from, either of type matrix or graphNEL.
- `to`: Character giving class to coerce to. Either "matrix" or "graphNEL".
- `strict`: Strict object checking.

Details

In the functions `ftM2adjM` and `ftM2graphNEL`, a from/to matrix `ft` is converted into an adjacency matrix or a graphNEL object respectively. In `ft`, the first column represents the from nodes and the second column the to nodes.

To have unconnected nodes, use the `V` argument (see below). The `edgemode` parameter can be used to specify if the desired output is a directed or undirected graph.

The same edge must not occur twice in the from/to matrix. If edgemode is undirected, the edge (u,v) and (v,u) must only be specified once.
Coercions between matrix and graph representations

W is an optional vector of edge weights. The order of the edge weights in the vector should correspond to the order of the edges recorded in ft. If it is not specified, edge weights of 1 are assigned by default.

V is an optional vector of node names. All elements of ft must be contained in V, but not all names in V need to be contained in ft. If V is not specified, it is set to all nodes represented in ft. Specifying V is most useful for creating a graph that includes nodes with degree 0.

aM is an affiliation matrix as frequently used in social networks analysis. The rows of aM represent actors, and the columns represent events. An entry of "1" in the ith row and jth column represents affiliation of the ith actor with the jth event. Weighted entries may also be used. aM2bpG returns a graphNEL object with nodes consisting of the set of actors and events, and directed (possibly weighted) edges from the actors to their corresponding events. If plotted using Rgraphviz and the dot layout, the bipartite structure of the graph returned by aM2bpG should be evident.

An adjacency matrix can be coerced into a graphNEL using the as method. If the matrix is a symmetric matrix, then the resulting graph will be undirected, otherwise it will be directed.

Value

For ftM2graphNEL and aM2bpG, an object of class graphNEL. For ftM2adjM, a matrix (the adjacency matrix representation).

Author(s)

Denise Scholtens, Wolfgang Huber

Examples

```r
## From-To matrix
From <- c("A","A","C","C")
To <- c("B","C","B","D")
L <- cbind(From,To)
W <- 1:4
M1 <- ftM2adjM(L, W, edgemode="directed")
M2 <- ftM2adjM(L, W, edgemode="undirected")
stopifnot(all(M1+t(M1)==M2))

G1 <- ftM2graphNEL(L, W, edgemode="directed")
G2 <- ftM2graphNEL(L, W, edgemode="undirected")

## Adjacency matrix

From <- matrix(rnorm(100), nrow=10, ncol=10)
From <- (From+t(From)) > pi/4
rownames(From) <- colnames(From) <- LETTERS[1:10]

To <- as(From,"graphNEL")
Back <- as(To,"matrix")

stopifnot(all(From == Back))
```
combineNodes

Description

A function to combine, or collapse, a specified set of nodes in a graph.

Usage

```r
combineNodes(nodes, graph, newName, ...)  
## S4 method for signature 'character,graphNEL,character'  
combineNodes(nodes, graph, newName, collapseFunction=sum)
```

Arguments

- `nodes`: A set of nodes that are to be collapsed.
- `graph`: The graph containing the nodes.
- `newName`: The name for the new, collapsed node.
- `collapseFunction`: Function or character giving the name of a function used to collapse the edge weights after combining nodes. The default is to sum up the weights, but mean would be a useful alternative.
- `...`: Additional arguments for the generic.

Details

The nodes specified are reduced to a single new node with label given by `newName`. The in and out edges of the set of nodes are all made into in and out edges for the new node.

Value

An new instance of a graph of the same class as `graph` is returned. This new graph has the specified nodes reduced to a single node.

Author(s)

R. Gentleman

See Also

`inEdges`, `addNode`
Examples

\begin{verbatim}
V <- LETTERS[1:4]
edL1 <- vector("list", length=4)
names(edL1) <- V
for(i in 1:4)
    edL1[[i]] <- list(edges=c(2,1,4,3)[i], weights=sqrt(i))
gR <- graphNEL(nodes=V, edgeL=edL1, edgemode="directed")
gR <- addNode("M", gR)
gR <- addEdge("M", "A", gR, 1)
gR <- addEdge("B", "D", gR, 1)
gX <- combineNodes(c("B","D"), gR, "X")

gR <- addNode("K", gR)
gR <- addEdge(c("K","K"), c("D","B"), gR, c(5,3))
edgeWeights(combineNodes(c("B","D"), gR, "X"))$K
edgeWeights(combineNodes(c("B","D"), gR, "X", mean))$K
\end{verbatim}

DFS

Depth First Search

Description

This function implements algorithm 4.2.1 of Gross and Yellen. The input is a graph and a node to start from. It returns a standard vertex labeling of graph. This is a vector with elements corresponding to the nodes of graph and with values that correspond to point in the depth first search the node is visited.

Usage

DFS(object, node, checkConn=TRUE)

Arguments

object An instance of the graph class.
node A character indicating the starting node.
checkConn A logical indicating whether the connectivity of the graph should be checked.

Details

This function implements algorithm 4.2.1 of Gross and Yellen. Specific details are given there.

It requires that the graph be connected. By default, this is checked, but since the checking can be expensive it is optional.

A faster and mostly likely better implementation of depth first searching is given by dfs in the RBGL package.
Value

A vector with names given by the nodes of graph whose values are 0 to one less than the number of nodes. These indices indicate the point at which the node will be visited.

Author(s)

R. Gentleman

References

*Graph Theory and its Applications*, J. Gross and J. Yellen.

See Also

boundary

Examples

```r
RNGkind("Mersenne-Twister")
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=.3)
RNGkind()
DFS(g1, "a")
```

---

**distGraph-class Class “distGraph”**

Description

A class definition for graphs that are based on distances.

Objects from the Class

Objects can be created by calls of the form `new("distGraph", ...).`

Slots

- **Dist**: Object of class "dist" that forms the basis for the edge weights used in the `distGraph`.

Extends

Class "graph", directly.
Methods

- **show** signature(object = "distGraph"): a print method
- **Dist** signature(object = "distGraph"): return the dist object.
- **adj** signature(object = "distGraph"): find the nodes adjacent to the supplied node.
- **nodes** signature(object = "distGraph"): return the nodes in the graph.
- **numNodes** signature(object = "distGraph"): return the number of nodes.
- **threshold** signature(object = "distGraph", k, value): set all distances that are larger than the supplied threshold, k, to the supplied value. The default is value is zero (and so is appropriate for similarities, rather than distances).
- **initialize** signature(object = "distGraph"): initialize a distGraph instance.
- **edgeWeights** Return a list of edge weights in a list format similar to the edges method.
- **edgeL** signature(graph = "distGraph"): A method for obtaining the edge list.

Author(s)

R. Gentleman

References

Shamir's paper and Butte et al

See Also

- `graph-class`
- `clusterGraph-class`

Examples

```r
set.seed(123)
x <- rnorm(26)
names(x) <- letters
library(stats)
d1 <- dist(x)
g1 <- new("distGraph", Dist=d1)
```

```
duplicatedEdges     duplicatedEdges
---------------------   ---------------------
```

Description

A multigraph is a graph where edges between nodes can be represented several times. For some algorithms this causes problems. *duplicatedEdges* tests an instance of the graphNEL class to see if it has duplicated edges and returns TRUE if it does and FALSE otherwise.

Usage

`duplicatedEdges(graph)`
edgeData-methods

Arguments

graph An instance of the class graphNEL

Details

It would be nice to handle other types of graphs.

Value

A logical, either TRUE if the graph has duplicated edges or FALSE if not.

Author(s)

R. Gentleman

See Also

connComp, ugraph

Examples

##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,

---

edgeData-methods Get and set attributes for the edges of a graph object

Description

Attributes of the edges of a graph can be accessed using edgeData. The attributes must be defined using edgeDataDefaults. You can omit the from or to argument to retrieve attribute values for all edges to (respectively, from) a given node.

Usage

dataframe(self, from, to, attr)
dataframe(self, from, to, attr) <- value

Arguments

self A graph-class instance
from A character vector of node names
to A character vector of node names
attr A character vector of length one specifying the name of a node attribute
value An R object to store as the attribute value
edgeDataDefaults-methods

*Get and set default attributes for the edges of a graph*

**Description**

Set default values for attributes associated with the edges of a graph.

**Usage**

```r
edgeDataDefaults(self, attr)
edgeDataDefaults(self, attr) <- value
```

**Arguments**

- `self`: A `graph-class` instance.
- `attr`: A character vector of length one giving the name of the attribute.
- `value`: An R class to use as the default value for the specified attribute.

---

**edgeMatrix**

*Compute an Edge Matrix or weight vector for a Graph*

**Description**

For our purposes an edge matrix is a matrix with two rows and as many columns as there are edges. The entries in the first row are the index of the node the edge is from, those in the second row indicate the node the edge is to.

If the graph is “undirected” then the duplicates option can be used to indicate whether reciprocal edges are wanted. The default is to leave them out. In this case the notions of from and to are not relevant.

**Usage**

```r
edgeMatrix(object, duplicates=FALSE)
eWV(g, eM, sep = ifelse(edgemode(g) == "directed", 
                   "->", 
                   "--"), useNNames=FALSE)
pathWeights(g, p, eM=NULL)
```
Arguments

- **object**: An object that inherits from `graph`.
- **g**: An object that inherits from `graph`.
- **duplicates**: Whether or not duplicate edges should be produced for “undirected” graphs.
- **eM**: An edge matrix.
- **sep**: A character string to concatenate node labels in the edge label.
- **useNNames**: A logical; if TRUE, node names are used in the edge label; if FALSE, node indices are used.
- **p**: A vector of node names constituting a path in graph `g`.

Details

Implementations for `graphNEL`, `clusterGraph` and `distGraph` are available.

Value

`edgeMatrix` returns a matrix with two rows, `from` and `to`, and as many columns as there are edges. Entries indicate the index in the node vector that corresponds to the appropriate end of the edge. `eWV` uses the edge matrix to create an annotated vector of edge weights. `pathWeights` returns an annotated vector of edge weights for a specified path in a graph.

Note

A path through an undirected graph may have several representations as a named vector of edges. Thus in the example, when the weights for path b-a-i are requested, the result is the pair of weights for edges a-b and a-i, as these are the edge labels computed for graph `g1`.

Author(s)

R. Gentleman

See Also

- `edges`

Examples

```r
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=0.3)
edgeMatrix(g1)
g2 <- new("clusterGraph", clusters=list(a=c(1,2,3), b=c(4,5,6)))
em2 <- edgeMatrix(g2)
eWV(g1, edgeMatrix(g1))
eWV(g1, edgeMatrix(g1), useNNames=TRUE)
pathWeights(g1, c("b", "a", "i"))
```
edgeSets

Description
C57BL/6J and C3H/HeJ mouse strains exhibit different cardiovascular and metabolic phenotypes on the hyperlipidemic apolipoprotein E (Apoe) null background. The interaction data for the genes from adipose, brain, liver and muscle tissue samples from male and female mice are included as a list of `data.frame`s. Each `data.frame` contains information for the from-gene, to-gene and the strength of interaction (weight) for each of the tissues studied.

Usage
```
data(esetsFemale)
data(esetsMale)
```

Source
Sage Commons Repository [http://sagebase.org/commons/dataset1.php#UCLA1](http://sagebase.org/commons/dataset1.php#UCLA1)

Examples
```
data(esetsFemale)
data(esetsMale)
```

edgeWeights

Description
A generic function that returns the edge weights of a graph. If `index` is specified, only the weights for the edges from the specified nodes are returned. The user can control which edge attribute is interpreted as the weight, see the Details section.

Usage
```
edgeWeights(object, index, ..., attr = "weight", default = 1, type.checker = is.numeric)
```
**edgeWeights**

**Arguments**

- **object**: A graph, any object that inherits from the graph class.
- **index**: If supplied, a character or numeric vector of node names or indices.
- **attr**: The name of the edge attribute to use as a weight. You can view the list of defined edge attributes and their default values using edgeDataDefaults. The default attribute name is "weight", see the Details section.
- **default**: The value to use if object has no edge attribute named by the value of attr. The default is the value 1 (double).
- **type.checker**: A function that will be used to check that the edge weights are of the correct type. This function should return TRUE if the input vector is of the right type and FALSE otherwise. The default is to check for numeric edge weights using is.numeric. If no type checking is desired, specify NULL.

**Details**

If index is supplied, then edge weights from these nodes to all adjacent nodes are returned. If index is not supplied, then the edge weights for all nodes are returned. The value for nodes without any outgoing edges will be a zero-length vector of the appropriate mode.

The edgeWeights method is a convenience wrapper around edgeData, the general-purpose way to access edge attribute information for a graph instance. In general, edge attributes can be arbitrary R objects. However, for edgeWeights to make sense, the values must be vectors of length not more than one.

By default, edgeWeights looks for an edge attribute with name "weight" and, if found, uses these values to construct the edge weight list. You can make use of attributes stored under a different name by providing a value for the attr argument. For example, if object is a graph instance with an edge attribute named "WTS", then the call edgeWeights(object, attr="WTS") will attempt to use those values.

The function specified by type.checker will be given a vector of edge weights; if the return value is not TRUE, then an error will be signaled indicating that the edge weights in the graph are not of the expected type. Type checking is skipped if type.checker is NULL.

If the graph instance does not have an edge attribute with name given by the value of the attr argument, default will be used as the weight for all edges. Note that if there is an attribute named by attr, then its default value will be used for edges not specifically customized. See edgeData and edgeDataDefaults for more information.

Because of their position after the ..., no partial matching is performed for the arguments attr, default, and type.checker.

**Value**

A named list of named edge weight vectors. The names on the list are the names of the nodes specified by index, or all nodes if index was not provided. The names on the weight vectors are node names to identify the edge to which the weight belongs.
fromGXL-methods

Methods for GXL manipulations in package graph

Description

GXL [http://www.gupro.de/GXL](http://www.gupro.de/GXL) is "an XML sublanguage designed to be a standard exchange format for graphs". This document describes tools in the graph package for importing GXL data to R and for writing graph data out as GXL.

Value

- **fromGXL** currently returns a graphNEL when possible. This function is based on `xmlEventParse` with handlers defined in the function `NELhandler`. The dump() element of this handler should emit information on all children of nodes and edges; the `asGraphNEL()` element will return a `graphNEL` object with weights if child `<attr>` with name attribute "weights" is present for each edge element.
- **toGXL** for an input of class "graphNEL", returns an object of class c("XMLInternalDOM", "XMLOutputStream"); see the example for how to convert this to a text stream encoding XML.
- **dumpGXL** returns an R list with all the node, edge, and named attribute information specified in the GXL stream.
- **validateGXL** returns silently (invisibly returns the parsed tree) for a DTD-compliant stream, or is otherwise very noisy.

Examples

```r
V <- LETTERS[1:4]
edL2 <- vector("list", length=4)
names(edL2) <- V
for(i in 1:4)
edL2[[i]] <- list(edges=c(2,1,2,1)[i], weights=sqrt(i))
gR2 <- graphNEL(nodes=V, edgeL=edL2, edgemode="directed")
edgeWeights(gR2, "C")
edgeWeights(gR2)
edgeWeights(gR2, attr="foo", default=5)
edgeData(gR2, attr="weight")
edgeData(gR2, from="C", attr="weight")
```
fromGXL-methods

Methods

fromGXL `con = connection`: returns a graphNEL based on a parsing of the GXL stream on the connection

dumpGXL `con = connection`: returns an R list based on a parsing of the GXL stream on the connection

validateGXL `con = connection`: checks the GXL stream against its DTD

toGXL `object = graphNEL`: creates an XMLInternalDOM representing the graph in GXL

Note

At present, toGXL does not return a validating GXL stream because XML package does not properly handle the dtd and namespaces arguments to xmlTree. This is being repaired. To fix the stream, add `<!DOCTYPE gxl SYSTEM "http://www.gupro.de/GXL/gxl-1.0.1.dtd">` as second record in the output.

Some structures in a graphNEL and some tags in GXL may not be handled at this time.

Author(s)

Vince Carey <stvjc@channing.harvard.edu>

Examples

```r
sf <- file(system.file("GXL/simpleExample.gxl", package="graph"))
show(fromGXL(sf))
print(dumpGXL(sf))
close(sf)
#validateGXL(sf)
# bad <- file(system.file("GXL/c2.gxl", package="graph"))
# here's how you can check if the GXL is well-formed, if
# you have a libxml2-based version of R XML package
#
# try( validateGXL(bad) )
#
gR <- graphNEL(nodes=letters[1:4], edgeL=list(a=list(edges=4), b=list(edges=3), c=list(edges=c(2,1)), d=list(edges=1)), edgemode="directed")
# following requires that you are using XML bound with recent libxml2
#
# an <- as.numeric
# if (an(libxmlVersion()$major)>=2 & an(libxmlVersion()$minor)>=4)
# since toGXL returns an XML object, we need to attach the XML package.
# library("XML")
cat(saveXML(toGXL(gR)$value()))
wtd <- file(system.file("GXL/kmstEx.gxl", package="graph"))
wtdg <- fromGXL(wtd)
close(wtd)
print(edgeWeights(wtdg))
```
# graph-class

## Description

A virtual class that all graph classes should extend.

## Details

- **degree** returns either a named vector (names correspond to the nodes in the graph) containing the degree for undirected graphs or a list with two components, `inDegree` and `outDegree` for directed graphs.
- **connComp** returns a list of the connected components. Each element of this list contains the labels of all nodes in that component.

For a *directed graph* or *digraph* the underlying graph is the graph that results from removing all direction from the edges. This can be achieved using the function `ugraph`. A *weakly connected* component of a *digraph* is one that is a connected component of the underlying graph. This is the default for `connComp`. A *digraph* is *strongly connected* if every two vertices are mutually reachable. A *strongly connected* component of a *digraph*, $D$, is a maximal *strongly connected* subdigraph of $D$. See the RBGL package for an implementation of Trajan’s algorithm to find *strongly connected* components (`strongComp`).

In the `graph` implementation of `connComp` weak connectivity is used. If the argument to `connComp` is a directed graph then `ugraph` is called to create the underlying undirected graph and that is used to compute connected components. Users who want different behavior are encouraged to use RBGL.

## Objects from the Class

A virtual Class: No objects may be created from it.

## Slots

- **edgeData**: An `attrData` instance for edge attributes.
- **nodeData**: An `attrData` instance for node attributes.
- **graphData**: A list for graph-level attributes. Only mandatory list item is `edgemode` which indicates whether edges are "directed" or "undirected".
- **renderInfo**: A list of graph rendering information.

## Methods

- **nodes** return a character vector containing the names of the nodes of the graph
- **nodes<-** rename the nodes of the graph
- **show** signature(object = "graph"): A print method for the graph.
- **acc** signature(object = "graph"): find all nodes accessible from the specified node.
complement signature(x = "graph"): compute the complement of the supplied graph. The complement is defined with respect to the complete graph on the nodes in x. Currently this returns an object of class graphNEL.

cconnComp signature(object = "graph"): find the connected components of a graph.

degree signature(object = "graph"): find the degree of a node (number of coincident edges).

degree signature(object = "MultiGraph"): find the degree of a node (number of coincident edges).

dfs signature(object = "graph"): execute a depth first search on a graph starting with the specified node.

edges signature(object="graph", which="character"): return the edges indicated by which. which can be missing in which case all edges are returned or it can be a character vector with the node labels indicating the nodes whose edge lists are wanted.

degreeDataDefaults Get and set default attributes for the edges in the graph.

degree Get and set attributes for edges in the graph

edgemode signature(object="graph"): return the edgemode for the graph. Currently this can be either directed or undirected.

edgemode<- signature(object="graph", value="character"): set the edgemode for the graph. Currently this can be either directed or undirected.

edgeWeights Return a list of edge weights in a list format similar to the edges method.

intersection signature(x = "graph", y = "graph"): compute the intersection of the two supplied graphs. They must have identical nodes. Currently this returns an object of class graphNEL. With edge weights of 1 for any matching edge.

isAdjacent signature(from="character", to="character"): Determine if edges exists between nodes.

isConnected signature(object = "graph"): A boolean that details if a graph is fully connected or not.

isDirected Return TRUE if the graph object has directed edges and FALSE otherwise.

join signature(x = "graph", y = "graph"): returns the joining of two graphs. Nodes which are shared by both graphs will have their edges merged. Note that edgeWeights for the resulting graph are all set to 1. Users wishing to preserve weights in a join operation must perform addEdge operations on the resulting graph to restore weights.

nodes<- A generic function that allows different implementations of the graph class to reset the node labels

nodeDataDefaults Get/set default attributes for nodes in the graph.

nodeData Get/set attributes for nodes in the graph.

numEdges signature(object = "graph"): compute the number of edges in a graph.

numNodes signature(object = "graph"): compute the number of nodes in a graph.

plot Please see the help page for the plot, graph-method method in the Rgraphviz package

union signature(x = "graph", y = "graph"): compute the union of the two supplied graphs. They must have identical nodes. Currently this returns an object of class graphNEL.

degreeNames signature(object = "graph"): Returns a vector of the edge names for this graph, using the format tail\~head, where head is the name of the tail node and head is the name of the head node.

updateGraph signature(object = "graph"): Updates old instances of graph objects.
Author(s)

R. Gentleman and E. Whalen.

References

Graph Theory and its Applications, J. Gross and J. Yellen.

See Also

graphNEL-class, graphAM-class, distGraph-class.

Examples

```r
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p= 0.3)
numEdges(g1)
edgeNames(g1)
edges(g1)
edges(g1, c("a","d")) # those incident to 'a' or 'd'
```

Description

The functions or variables listed here are no longer part of the graph package.

Usage

```r
buildRepDepGraph()
pkgInstOrder()
ugraphOld()
```

See Also

Defunct
Graphical parameters and other settings

Description

Functions providing an interface to persistent graphical parameters and other settings used in the package.

Usage

graph.par(...)
graph.par.get(name)

Arguments

... either character strings naming parameters whose values are to be retrieved, or named arguments giving values that are to be set.
name character string, giving a valid parameter name.

Details

graph.par works sort of like par, but the details are yet to be decided.
graph.par.get(name) is equivalent to graph.par(name)[[1]]

Value

In query mode, when no parameters are being set, graph.par returns a list containing the current values of the requested parameters. When called with no arguments, it returns a list with all parameters. When a parameter is set, the return value is a list containing previous values of these parameters.

Author(s)

Deepayan Sarkar, <deepayan.sarkar@r-project.org>

See Also

par
graph2SparseM  

Coercion methods between graphs and sparse matrices

Description

These functions provide coercions between objects that inherit from the graph class to sparse matrices from the SparseM package.

Usage

\[
\text{graph2SparseM}(g, \text{useweights}=\text{FALSE}) \\
\text{sparseM2Graph}(sM, \text{nodeNames}, \text{edgemode}=c(\text{"directed"}, \text{"undirected"}))
\]

Arguments

- \text{g}\quad\text{An instance of the graph class.}
- \text{useweights}\quad\text{A logical value indicating whether to use the edge weights in the graph as values in the sparse matrix.}
- \text{sM}\quad\text{A sparse matrix.}
- \text{nodeNames}\quad\text{A character vector of the node names.}
- \text{edgemode}\quad\text{Specifies whether the graph to be created should have directed (default) or undirected edges. If undirected, the input matrix sM must be symmetric.}

Details

A very simple coercion from one representation to another.

Currently it is presumed that the matrix is square. For other graph formats, such as bipartite graphs, some improvements will be needed; patches are welcome.

Value

- graph2SparseM takes as input an instance of a subclass of the graph class and returns a sparse matrix.
- sparseM2Graph takes a sparse matrix as input and returns an instance of the graphNEL class. By default, the graphNEL returned will have directed edges.

Author(s)

R. Gentleman

See Also

\texttt{graph-class}, \texttt{graphNEL-class}, and for other conversions, \texttt{aM2bG} and \texttt{ftM2adjM}
Examples

```r
set.seed(123)
g1 <- randomGraph(letters[1:10], 1:4, p=.3)
s1 <- graph2SparseM(g1, useweights=TRUE)
g2 <- sparseM2Graph(s1, letters[1:10], edgemode="undirected")
## consistency check
stopifnot(all.equal(g1, g2))
```

---

**Description**

A graph class where node and edge information is represented as an adjacency matrix. The adjacency matrix is square and element `adjMat[i, j]` is one if there is an edge from node `i` to node `j` and zero otherwise.

**Details**

The non-zero matrix values can be used to initialize an edge attribute. If this is desired, use the `values` argument in the call to `new` and provide a list with a single named element. The name determines the attributes and the value provides the default value for that attribute.

**Objects from the Class**

Objects can be created by calls of the form `graphAM(adjMat, edgemode, values)`.

**Slots**

- `adjMat`: An adjacency "matrix" describing the graph structure. The `colnames` of the matrix will be used as node names for the graph if present.
- `edgeData`: Storage for edge attributes.
- `nodeData`: Storage for node attributes.

**Extends**

Class "graph", directly.

**Constructor**

`graphAM(adjMat=matrix(integer(), 0, 0), edgemode='undirected', values=NA)` creates a `graphAM` instance.

- `adjMat` An integer matrix specifying which nodes have edges between them.
- `edgemode` Either "directed" or "undirected".
values A named list of length 1, used (rather obscurely) to specify that non-zero adjMat values initialize an edge attribute. The name of the single element in that list becomes the name of that attribute, with the specified default value. This default value is, however, never used: the specified edge attribute always has the value contained in the adjacency matrix, which is traditionally 1, but can be any positive number.

Methods

```r
addEdge signature(from = "character", to = "character", graph = "graphAM", weights = "missing"): ...
addNode signature(object = "graphAM", nodes = "character"): ...
clearNode signature(node = "character", object = "graphAM"): ...
coerce signature(from = "graphAM", to = "graphNEL"): ...
coerce signature(from = "graphAM", to = "graphBAM"): ...
coerce signature(from = "graphAM", to = "matrix"): In converting to a matrix, if an edge attribute named "weight" is defined, the non-zero elements of the matrix will contain the corresponding attribute value. For more flexible matrix conversion, see toMatrix.
coerce signature(from = "matrix", to = "graphAM"): This coerce method exists for symmetry.
   In most cases, creating a new graphAM instance using new gives one more control over the resulting graph.
edges signature(object = "graphAM"): ...
initialize signature(.Object = "graphAM"): ...
inEdges signature(node = "character", object = "graphNEL"): Return the incoming edges for the specified nodes. See inEdges.
isAdjacent signature(object = "graphAM", from = "character", to = "character"): ...
nodes<- signature(object = "graphAM", value = "character"): ...
nodes signature(object = "graphAM"): ...
numEdges signature(graph = "graphAM"): ...
numNodes signature(object = "graphAM"): ...
removeEdge signature(from = "character", to = "character", graph = "graphAM"): ...
removeNode signature(node = "character", object = "graphAM"): ...
```

Author(s)

Seth Falcon

See Also

```
graph-class, graphNEL-class
```
Examples

```r
mat <- rbind(c(0, 0, 1, 1),
             c(0, 0, 1, 1),
             c(1, 1, 0, 1),
             c(1, 1, 1, 0))
rownames(mat) <- colnames(mat) <- letters[1:4]
g1 <- graphAM(adjMat=mat)
stopifnot(identical(mat, as(g1, "matrix")), validObject(g1))

## now with weights:
mat[1,3] <- mat[3,1] <- 10
gw <- graphAM(adjMat=mat, values=list(weight=1))

## consistency check:
stopifnot(identical(mat, as(gw, "matrix")),
          validObject(gw),
          identical(gw, as(as(gw, "graphNEL"), "graphAM")))
```

Description

The graphBAM class represents a graph as an adjacency matrix. The adjacency matrix is stored as a bit array using a raw vector to reduce the memory footprint and speed operations like graphIntersection. This class is EXPERIMENTAL and its API is subject to change.

Usage

```r
graphBAM(df, nodes=NULL, edgemode="undirected", ignore_dup_edges = FALSE)
```

Arguments

- `df` A data.frame with three columns: "from", "to" and "weight". Columns "from" and "to" can be either factors or character vectors. Each row of `df` describes an edge in the resulting graph. The "weight" column must be numeric.
- `nodes` A character vector of node labels. Use this to add degree zero nodes to the graph. If `NULL`, the set of nodes found in `from` and `to` will be used.
- `edgemode` A string, one of "directed" or "undirected".
- `ignore_dup_edges` If `FALSE` (default), specifying duplicate edges in the input is an error. When set to `TRUE` duplicate edges are ignored. Edge weight values are ignored when determining duplicates. This is most useful for graph import and conversion.

Constructors

The `GraphBAM` function is used to create new `graphBAM` instances. Edges are specified in a `data.frame`. For undirected graphs, reciprocal edges should not be includes unless `ignore_dup_edges` is `TRUE`. 
Extends
Class "graph", directly.

Methods for graphBAM objects

addEdge(from, to, graph, weights) Return a new graphBAM object with the specified edge(s) added. The from and to arguments must either be the same length or one of them must be of length one. Each time an edge is added, the entire graph is copied. For the purpose of building a graph it will often be more efficient to build up the list of edges and call GraphBAM.

addNode(node, object) Return a new graphBAM object with the specified node(s) added.

clearNode(node, object) This operation is not currently supported.

edges(object, which) Returns an adjacency list representation of the graph. The list will have an entry for each node with a vector of adjacent node labels or character(0). For undirected graphs, edges returns the reciprocal edges. The optional argument which can be a character vector of node labels. When present, only entries for the specified nodes will be returned.

inEdges(node, object) (Not yet supported) Similar to the edges function, but the adjacency list maps nodes that have an edge to the given node instead of from the given node.

isAdjacent(object, from, to) Returns a logical vector indicating whether there is an edge corresponding to the elements in from and to. These vectors must have the same length, unless one has length one.

nodes(object) Return the node labels for the graph

numEdges(object) Returns the number of edges in the graph.

numNodes(object) Returns the number of nodes in the graph.

removeEdge(from, to, graph) Return a new graphBAM object with the specified edges removed. The from and to arguments must be the same length unless one of them has length one.

removeNode(node, object) Returns a new graphBAM object with the specified node removed. Node and edge attributes corresponding to that node are also removed.

edgeData(self, from, to, attr) Access edge attributes. See help for edgeData.

edgeDataDefaults(self, attr) Access edge data default attributes.

nodeDataDefaults(self, attr) Access node data default attributes.

edgeWeights(object, index) Return the edge weights for the graph in adjacency list format. The optional argument index specified a character vector of nodes. In this case, only the weights for the specified nodes will be returned.

extractFromTo(g) Returns a data frame with column names "from", "to", and "weight" corresponding to the connected nodes in the graphBAM object.

graphIntersect(x, y, nodeFun, edgeFun) When given two graphBAM objects, graphIntersect returns a new graphBAM containing the nodes and edges in common between the two graphs. Both x and y should either be directed or undirected. The intersection is computed by first finding the intersection of the node sets, obtaining the resulting subgraphs, and finding the intersection of the resulting edge sets. Node/Edge attributes that are equal are carried over to the result. Non equal edge/node attributes will result in the corresponding attribute being set to NA. The user has the option of providing a named list of functions corresponding to the names of the edge attributes for resolving conflicting edge attributes. For resolving any of the conflicting node attributes the user has the option of providing a named list of functions corresponding to the node attribute names.
graphUnion(x, y, nodeFun, edgeFun) When given two graphBAM objects, graphUnion returns a new graphBAM containing the union of nodes and edges between the two graphs. The union is computed by first finding the union of the nodesets. Both x and y should be either directed or undirected. Node/Edge attributes that are equal are carried over to the result. Non equal edge/node attributes will result in the corresponding attribute being set to NA. The user has the option of providing a named list of functions corresponding to the names of the edge attributes for resolving conflicting edge attributes. For resolving any of the conflicting node attributes the user has the option of providing a named list of functions corresponding to the node attribute names.

edgemode(object) <- value Set the edgemode for the graph ("directed" or "undirected"). If the specified edgemode is the same, the object is returned without changes. Otherwise, a directed graph is converted to an undirected graph via ugraph and an undirected graph is returned such that each edge is interpreted as two edges, one in each direction.

ugraph(graph) Return an undirected version of the current graph. Conceptually, the arrows of a graph’s directed edges are removed.

nodes(object) <- value Replacement of a graphBAM object’s node labels is currently not supported. An error is raised if this method is called.

Coercion

graphBAM objects can be coerced to graphAM, graphNEL, and matrix instances via as(g, CLASS).

Author(s)

N. Gopalakrishnan, S. Falcon

Examples

```r
f <- c("a", "a", "b", "c", "d")
t <- c("b", "c", "c", "d", "a")
weight <- c(2.3, 2.3, 4.3, 1.0, 3.0)
df <- data.frame(from=f, to=t, weight= weight, stringsAsFactors = TRUE)
g <- graphBAM(df)
nd <- nodes(g)
nodeDataDefaults(g, attr ="color") <- "green"
nodeData(g,n=c("b", "c"), attr ="color") <- "red"
w1 <- edgeWeights(g)
w2 <- edgeWeights(g,"a")
w3 <- edgeWeights(g,1)
d1 <- edges(g)
d2 <- edges(g,c("a", "b"))
e1 <- edgeData(g)
e2 <- edgeData(g, "a", "c",attr="weight")
e3 <- edgeData(g)
id <- isDirected(g)
sg <- subGraph(c("a","c","d"), g)
fT <- extractFromTo(g)
am <- as(g,"graphAM")
nl <- as(g,"graphNEL")
mT <- as(g,"matrix")
```
k <- graphIntersect(g,g)
k <- graphUnion(g,g)
e <- removeEdgesByWeight(g, lessThan=3.0)
f <- removeNode("a", g)
g

graphExamples  

A List Of Example Graphs

**Description**

This data set contains a list of example graphNEL objects, which can then be used for plotting.

**Usage**

data(graphExamples)

**Source**

Various sources, primarily from `randomGraph` and `randomEGraph`

**Examples**

data(graphExamples)
a <- graphExamples[[1]]
a

graphNEL-class  

Class "graphNEL"

**Description**

This is a class of graphs that are represented in terms of nodes and an edge list. This is a suitable representation for a graph with a large number of nodes and relatively few edges.

**Details**

The `graphNEL` class provides a very general structure for representing graphs. It will be reasonably efficient for lists with relatively more nodes than edges. Although this representation can support multi-edges, such support is not implemented and instances of `graphNEL` are assumed to be simple graphs with at most one edge between any pair of nodes.

The `edgeL` is a named list of the same length as the node vector. The names are the names of the nodes. Each element of `edgeL` is itself a list. Each element of this (sub)list is a vector (all must be the same length) and each element represents an edge to another node. The sublist named `edges` holds index values into the node vector. And each such entry represents an edge from the node which has the same name as the component of `edgeL` to the node with index provided. Another
component that is often used is named weights. It represents edge weights. The user can specify any other edge attributes (such as types etc). They are responsible for any special handling that these might require.

For an undirected instance all edges are reciprocated (there is an edge from A to B and from B to A).

Note that the reason for using indices to represent the to end of a node is so that we can easily support permutation of the node labels as a way to generate randomizations of the graph.

Slots

nodes: Object of class "vector".

edgeL: Object of class "list". The edgeL must be the same length as nodes. The elements of this vector correspond to the same element in nodes. The elements are themselves lists. If the node has any edges then this list will have an element named edges. This will eventually change. Since edge weights are now stored in the edge attributes construct, we do not need the extra level of list.

Extends

Class "graph", directly.

Constructor

graphNEL(nodes=character(), edgeL=list(), edgemode='undirected') creates a graphNEL instance.

nodes A character vector of node labels.

edgel A named list either in the format returned by the edges method or a list of lists where each inner list has an element named edges and optionally an element named weights. If weights is present, it must be the same length as the edges element.

edgemode Either "directed" or "undirected".

Methods

adj signature(object = "graphNEL"): A method for finding nodes adjacent to the supplied node.

edgel signature(graph = "graphNEL"): A method for obtaining the edge list.

edgeWeights signature(object = "graphNEL"): A method for obtaining the edge weights.

edges signature(object = "graphNEL"): A method for obtaining the edges.

inEdges signature(node = "character", object = "graphNEL"): Return the incoming edges for the specified nodes. See inEdges.

nodes signature(object = "graphNEL"): A method for obtaining the nodes.

numNodes signature(object = "graphNEL"): A method for determining how many nodes are in the graph.

subGraph signature(snodes="character", graph = "graphNEL"): A method for obtaining the induced subgraph based on the set of supplied nodes and the supplied graph.

plot Please see the help page for plot.graphNEL in the Rgraphviz package
**graph2graphviz** signature(object = "graphNEL"): A method that will convert a graphNEL object into a matrix suitable for interaction with Rgraphviz. Not intended to be called directly. This function will insure that no NA's (or other undesired values) are in the graph, or created by coercion.

**nodes**<- signature(object="graphNEL", value="character"): A method for replacing the nodes in a graph object. It checks to be sure the values are the right length and unique.

**coerce** signature(from = "graphNEL", to = "graphAM"): Called via as, the method converts to an adjacency matrix representation. See **graphAM-class**.

**coerce** signature(from = "graphNEL", to = "graphBAM"): Called via as, the method converts to an bit array representation. See **graphBAM-class**.

**Author**

R. Gentleman

**See Also**

**graphAM-class, distGraph-class, clusterGraph-class**

**Examples**

```r
set.seed(123)
V <- LETTERS[1:4]
edL <- vector("list", length=4)
names(edL) <- V
for(i in 1:4)
edL[[i]] <- list(edges=5-i, weights=runif(1))
gR <- graphNEL(nodes=V, edgeL=edL)
edges(gR)
edgeWeights(gR)
```

---

**inEdges**

**Generic Method inEdges**

**Description**

Returns a list of all incoming edges for the specified nodes.

**Usage**

`inEdges(node, object)`

**Arguments**

- **node** character vector of node names
- **object** a graph object
Details

If no node argument is specified, `inEdges` returns the incoming edges for all nodes in the graph.
For an undirected graph, `inEdges` returns all edges for the specified nodes.

Value

A list with length matching the length of `node`. If `node` was missing, a list containing an element for each node in the graph.
Each list element contains a character vector of node names giving the nodes that have outgoing edges to the node given by the name of the list element.

Author(s)

R. Gentleman

See Also

`removeNode`, `clearNode`

Examples

```r
V <- LETTERS[1:4]
edL3 <- vector("list", length=4)
for(i in 1:4)
edL3[[i]] <- list(edges=(i%%4)+1, weights=i)
names(edL3) <- V
gR3 <- graphNEL(nodes=V, edgeL=edL3, "directed")
inEdges(c("A", "B"), gR3)
```
Details

The integrinMediatedCellAdhesion data set contains two objects:

The first is IMCAGraph, which is an object of class graph-NEL and represents the hsa04510 graph from KEGG.

The second is IMCAAttrs, which is a list of four elements. The first element, defAttrs corresponds to the attrs arguments of agopen and plot.graph. The second element is nodeAttrs which corresponds to the nodeAttrs argument in the same two functions from Rgraphviz. The third element, subGList corresponds to the subGList argument in those functions. Lastly, the fourth element, LocusLink provides a named list where the names are the nodes and the values are vectors of LocusLink ID values which correspond to those nodes.

The values from defAttrs, nodeAttrs and subGList in the IMCAAttrs list are part of an ongoing attempt by Bioconductor to provide the set of options to most accurately recreate the actual visual image of the pathway from the KEGG site using Rgraphviz. Users may try out their own combination of attributes and settings for their own needs, but these represent our own efforts at as closely recreating the image as possible.

Source


Examples

data(integrinMediatedCellAdhesion)
if (require("Rgraphviz") & interactive())
  plot(IMCAGraph, attrs=IMCAAttrs$defAttrs,
       nodeAttrs=IMCAAttrs$nodeAttrs, subGList=IMCAAttrs$subGList)

internal

Variables used for internal purposes

Description

The nullgraphID variable is used to store a default identifier. This should not be used by users.

Author(s)

Saikat DebRoy
isAdjacent-methods  Determine if nodes share an edge in a graph

Description

For a given subclass of graph-class, returns TRUE if the graph contains an edge from node specified by from to the node specified by to.

The appropriate logical vector will be returned as long as from and to have the same length and contain nodes in the graph object specified by object.

Usage

isAdjacent(object, from, to, ...)

Arguments

object  An instance of a subclass of graph-class.
from  A character vector of nodes in the graph.
to  A character vector of nodes in the graph
...  May be used by methods called on subclasses of graph

isDirected-methods  Determine if a graph has directed or undirected edges

Description

The edges of a graph-class object are either directed or undirected. This function returns TRUE if the edges are directed and FALSE otherwise.

Usage

isDirected(object)

Arguments

object  A graph-class instance
leaves

Find the leaves of a graph

Description

A leaf of an undirected graph is a node with degree equal to one. A leaf of a directed graph is defined with respect to in-degree or out-degree. The leaves of a directed graph with respect to in-degree (out-degree) are those nodes with in-degree (out-degree) equal to zero.

Usage

leaves(object, degree.dir)

Arguments

object  A graph object
degree.dir  One of "in" or "out". This argument is ignored when object is undirected and required otherwise. When degree.dir="in" (degree.dir="out"), nodes have no in coming (out going) edges will be returned.

Value

A character vector giving the node labels of the leaves.

Author(s)

Seth Falcon

Examples

data(graphExamples)
graphExamples[[1]]
leaves(graphExamples[[1]])

data(apopGraph)
leaves(apopGraph, "in")
leaves(apopGraph, "out")
List the Edges of a Graph

Description

A list where each element contains all edges between two nodes, regardless of orientation. The list has names which are node pairs, in lexicographic order, and elements all edges between those nodes.

Usage

listEdges(object, dropNULL=TRUE)

Arguments

object An instance of the graphNEL-class class.
dropNULL Should those node pairs with no edges be dropped from the returned list.

Details

The function is currently only implemented for graphs of the graphNEL-class class. The edges in the returned list are instances of the simpleEdge-class.

Value

A named list of simpleEdge-class objects.

Author(s)

R. Gentleman

See Also

simpleEdge-class, edges

Examples

set.seed(123)
V <- LETTERS[1:4]
edL <- vector("list", length=4)
names(edL) <- V
toE <- LETTERS[4:1]
for(i in 1:4)
  edL[[i]] <- list(edges=5-i, weights=runif(1))
gR <- graphNEL(nodes=V, edgeL=edL)
listEdges(gR)
MAPKsig

A graph encoding parts of the MAPK signaling pathway

Description

A graph encoding parts of the MAPK signaling pathway

Usage

data(MAPKsig)

Format

The format is: Formal class 'graphNEL' [package "graph"] with edgemode "directed".

Source

The KEGG pancreatic cancer pathway was visually inspected on 17 Sept 2007, and the subgraph associated with MAPK signaling was isolated. The HUGO names for each symbol in the KEGG visualization were obtained and checked for existence on hgu95av2. Some abbreviated terms for processes are also included as nodes.

Examples

data(MAPKsig)
if (require(Rgraphviz)) {
  nat = rep(FALSE, length(nodes(MAPKsig)))
  names(nat) = nodes(MAPKsig)
  plot(MAPKsig, nodeAttrs=list(fixedsize=nat))
}

mostEdges

Find the node in a graph with the greatest number of edges

Description

mostEdges finds the node that has the most edges in the graph. This is the node with the highest degree.

Usage

mostEdges(objGraph)

Arguments

objGraph the graph object
MultiGraph-class

**Value**

- **index** the index of the node with the most edges
- **id** the node value with the most edges; may be affy id, locus link id, or genename depending on the node type
- **maxLen** the number of edges for that node

**Author(s)**

Elizabeth Whalen

**See Also**

numEdges, aveNumEdges, numNoEdges

**Examples**

```r
set.seed(123)
g1 <- randomGraph(11:30, letters[20:26], p=.4)
mostEdges(g1)
```

---

**Description**

The MultiGraph class represents a single node set and a set of edge sets. Each edge set is either directed or undirected. We can think of an edge in a MultiGraph as a 4-tuple (from-node, to-node, edge-type, weight), where the edge-type field in the tuple identifies the edge set, the weight is a numeric value, and the order of the nodes only matters in the case of a directed edge set. Unlike some of the graph representations, self-loops are allowed (from-node == to-node).

There is support for arbitrary edge attributes which is primarily useful for rendering plots of MultiGraphs. These attributes are stored separately from the edge weights to facilitate efficient edge weight computation.

**Usage**

```r
MultiGraph(edgeSets, nodes = NULL, directed = TRUE, ignore_dup_edges = FALSE)
eweights(object, names.sep = NULL)
edgeSetIntersect0(g, edgeFun = NULL)
edgeSetIntersect0(g, edgeFun = NULL)
extractGraphAM(g, edgeSets)
extractGraphBAM(g, edgeSets)
```
Arguments

edgeSets A named list of data.frame objects each representing an edge set of the multigraph. Each data.frame must have three columns: "from", "to", and "weight". Columns "from" and "to" can be either factors or character vectors. The "weight" column must be numeric.

nodes A character vector of node labels. Nodes with zero degree can be included in a graph by specifying the node labels in nodes. The node set of the resulting multigraph is the union of the node labels found in edgeSets and nodes.

directed A logical vector indicating whether the edge sets specified in edgeSets represent directed edges. If this argument has length one, the value applies to all edge sets in edgeSets. Otherwise, this argument must have the same length as edgeSets, values are aligned by position.

object A MultiGraph instance

g A MultiGraph instance

names.sep The string to use as a separator between from and to node labels. If NULL no names will be attached to the returned vector.

ignore_dup_edges If FALSE (default), specifying duplicate edges in the input is an error. When set to TRUE duplicate edges are ignored. Edge weight values are ignored when determining duplicates. This is most useful for graph import and conversion.

edgeFun A user specified named list of functions to resolve edge attributes in a union or intersection operation

Constructors

MultiGraph

Methods

nodes Return the nodes of the multigraph.

numEdges Return an integer vector named by edge set containing edge counts for each edge set.

numNodes Return the number of nodes in the multigraph.

eweight Return a list named by edge set; each element is a numeric vector of edge weights for the corresponding edge set.

isDirected Return a logical vector named by the edge sets in object with a TRUE indicating a directed edge set and FALSE for undirected.

edges Returns a list named by edge set; for the edges in the MultiGraph

dgeNames Returns a list named by the edge set; for the names of the edges in the MultiGraph

extractFromTo Return a list named by the edge sets; each element is a data frame with column names from, to and weight corresponding to the connected nodes in the edge set.

subsetEdgeSets Return a new MultiGraph object representing the subset of edge sets from the original MultiGraph.

eextractGraphAM Return a named list of graphAM objects corresponding to the edge sets from the original MultiGraph.
extractGraphBAM  Return a named list of graphBAM objects corresponding to the edge sets from
the original MultiGraph.

ugraph Return a new MultiGraph object in which all edge sets have been converted to undirected
edge sets. This operation sets all edge weights to one and drops other edge attributes.

edgeSetIntersect0 Return a new MultiGraph object representing the intersection of edges across
all edge sets within g. The return value will have a single edge set if the edge sets in g are
disjoint. Otherwise, there will be a single edge set containing the shared edges. The node set
is preserved. Edge weights and edge attributes are transferred over to the output if they have
the same value, else user has the option of providing a function to resolve the conflict.

edgeSetUnion0 Return a new MultiGraph object representing the union of edges across all edge
sets within g. The node set is preserved. Edge weights and edge attributes are transferred over
to the output if they have the same value, else user has the option of providing a function to
resolve the conflict.

graphIntersect(x, y, nodeFun, edgeFun) When given two MultiGraph objects, graphIntersect
returns a new MultiGraph containing the nodes and edges in common between the two graphs.
The intersection is computed by first finding the intersection of the node sets, obtaining the
induced subgraphs, and finding the intersection of the resulting edge sets. The corresponding
named edgeSets in x and y should both be either directed or undirected.Node/Edge attributes
that are equal are carried over to the result. Non equal edge/node attributes will result in
the corresponding attribute being set to NA. The user has the option of providing a named
list(names of edgeSets) of list (names of edge attributes) of edge functions corresponding to the
names of the edge attributes for resolving conflicting edge attributes (edgeFun). For resolving
any of the conflicting node attributes, the user has the option of providing a named list of
functions corresponding to the node attribute names (nodeFun).

graphUnion(x, y, nodeFun, edgeFun) When given two MultiGraph objects, graphUnion re-
turns a new MultiGraph containing the union of nodes and edges between the two graphs.
The corresponding pairs of named edgeSets in x and y should both be either directed or undi-
rected. Non equal edge/node attributes will result in the corresponding attribute being set to
NA. The user has the option of providing a named list(names of edgeSets) of list (names of
dge attributes) of edge functions corresponding to the names of the edge attributes for resolv-
ing conflicting edge attributes (edgeFun). For resolving any of the conflicting node attributes,
the user has the option of providing a named list of functions corresponding to the node
attribute names (nodeFun).

dgeSets(object, ...) Returns the names of the edgeSets in the MultiGraph object as a char-
acter vector.

show Prints a short summary of a MultiGraph object

Author(s)

S. Falcon, Gopalakrishnan N

Examples

ft1 <- data.frame(from=c("a", "a", "a", "b", "b"),
     to=c("b", "c", "d", "a", "d"),
     weight=c(1, 3.1, 5.4, 1, 2.2),
stringsAsFactors = TRUE)
multigraph-class

Class "multigraph"

Description

A collection of classes to model multigraphs. These include the multigraph class as well as classes to contain edge sets.

Objects from the Class

Objects can be created from the multigraph class, the edgeSet class is virtual, and particular variants should be used.
Slots

These slots are for the multiGraph class.

- **nodes**: The names of the nodes.
- **edgeL**: A list of edge lists.
- **nodeData**: An instance of the attrData class.
- **graphData**: A list.

These slots are for the edgeSet class, or one of its subclasses.

- **edgeData**: An instance of the attrData class.
- **edgemode**: A character vector, one of directed, or undirected.
- **edgeL**: A list of the edges (graphNEL)
- **adjMat**: An adjacency matrix (graphAM)

Methods

- **show**: Print a multigraph.
- **isDirected**: A vector indicating which of the edgeSets is directed.
- **nodes**: Retrieve the node names
- **numNodes**: Return the number of nodes
- **edges**: Return either all edges, or a subset of them, depending on the arguments supplied.
- **numEdges**: Return a vector with the number of edges, for each edge set.

nodeData-methods

Get and set attributes for the nodes of a graph object

Description

Attributes of the nodes of a graph can be accessed using nodeData. The attributes must be defined using nodeDataDefaults. You can ommitt the n argument to retrieve attributes for all nodes in the graph. You can ommitt the attr argument to retrieve all attributes.

Usage

nodeData(self, n, attr)
nodeData(self, n, attr) <- value

Arguments

- **self**: A graph-class instance
- **n**: A character vector of node names
- **attr**: A character vector of length one specifying the name of a node attribute
- **value**: An R object to store as the attribute value
nodeDataDefaults-methods

Get and set default attributes for the nodes of a graph

Description

You can associate arbitrary attributes with the nodes of a graph. Use nodeDataDefaults to specify the set of attributes that describe nodes. Each attribute must have a default value. You can set the attribute for a particular node or set of nodes using nodeData.

Usage

nodeDataDefaults(self, attr)
nodeDataDefaults(self, attr) <- value

Arguments

self     A graph-class instance
attr     A character vector of length one giving the name of an attribute
value    An R object to set as the default value for the given attribute

numNoEdges

Calculate the number of nodes that have an edge list of NULL

Description

numNoEdges calculates the number of nodes that have an edge list of NULL (i.e. no edges).

Usage

numNoEdges(objGraph)

Arguments

objGraph  the graph object

Value

An integer representing the number of NULL edge lists in the graph.

Author(s)

Elizabeth Whalen
See Also

numEdges, aveNumEdges, mostEdges

Examples

```r
set.seed(999)
g1 <- randomEGraph(letters, .01)
numNoEdges(g1)
```

---

**pancrCaIni**

A graph encoding parts of the pancreatic cancer initiation pathway

**Description**

A graph encoding parts of the pancreatic cancer initiation pathway

**Usage**

data(pancrCaIni)

**Format**

The format is: Formal class 'graphNEL' [package "graph"] with edgemode "directed".

**Source**

The KEGG pancreatic cancer pathway was visually inspected on 17 Sept 2007 and a subgraph was isolated. The HUGO names for each symbol in the KEGG visualization were obtained and checked for existance on hgu95av2. Some abbreviated terms for processes are also included as nodes.

**Examples**

```r
data(pancrCaIni)
if (require(Rgraphviz)) {
  nat = rep(FALSE, length(nodes(pancrCaIni)))
names(nat) = nodes(pancrCaIni)
plot(pancrCaIni, nodeAttrs=list(fixedsize=nat))
}
```
randomEGraph  

Random Edge Graph

Description
A function to create random graphs according to a random edge model. The user supplies the set of nodes for the graph as V and either a probability, p, that is used for each edge or the number of edges, edges they want to have in the resulting graph.

Usage
randomEGraph(V, p, edges)

Arguments
- V: The nodes for the graph.
- p: The probability of an edge being selected.
- edges: The number of edges wanted.

Details
The user must specify the set of nodes and either a probability for edge selection or the number of edges wanted, but not both. Let nV denote the number of nodes. There are \( \binom{nV}{2} \) edges in the complete graph. If p is specified then a biased coin (probability of heads being p) is tossed for each edge and if it is heads that edge is selected. If edges is specified then that many edges are sampled without replacement from the set of possible edges.

Value
An object of class graphNEL-class that contains the nodes and edges.

Author(s)
R. Gentleman

See Also
randomGraph

Examples
```r
set.seed(123)
V <- letters[14:22]
g1 <- randomEGraph(V, .2)
g2 <- randomEGraph(V, edges=30)
```
randomGraph

Description

This function generates a random graph according to a model that involves a latent variable. The construction is to randomly assign members of the set M to the nodes, V. An edge is assigned between two elements of V when they both have the same element of M assigned to them. An object of class graphNEL is returned.

Usage

randomGraph(V, M, p, weights=TRUE)

Arguments

V
The nodes of the graph.

M
A set of values used to generate the graph.

p
A value between 0 and 1 that indicates the probability of selecting an element of M

weights
A logical indicating whether to use the number of shared elements of M as weights.

Details

The model is quite simple. To generate a graph, G, the user supplies the list of nodes, V and a set of values M which will be used to create the graph. For each node in V a logical vector with length equal to the length of M is generated. The probability of a TRUE at any position is determined by p. Once values from M have been assigned to each node in V the result is processed into a graph. This is done by creating an edge between any two elements of V that share an element of M (as chosen by the selection process).

The sizes of V and M and the values of p determine how dense the graph will be.

Value

An object of class graphNEL-class is returned.

Author(s)

R. Gentleman

See Also

randomEGraph, randomNodeGraph
Examples

```r
set.seed(123)
V <- letters[1:10]
M <- 1:4
g1 <- randomGraph(V, M, 0.2)
numEdges(g1) # 16, in this case
g1 edges (g1) # "from" - "to" since undirected
```

---

**randomNodeGraph**

*Generate Random Graph with Specified Degree Distribution*

**Description**

randomNodeGraph generates a random graph with the specified degree distribution. Self-loops are allowed. The resultant graph is directed (but can always be coerced to be undirected).

**Usage**

```r
randomNodeGraph(nodeDegree)
```

**Arguments**

- `nodeDegree` A named integer vector specifying the node degrees.

**Details**

The input vector must be named, the names are taken to be the names of the nodes. The sum must be even (there is a theorem that says we require that to construct a graph). Self-loops are allowed, although patches to the code that make this a switchable parameter would be welcome.

**Value**

An instance of the graphNEL class. The graph is directed.

**Author(s)**

R. Gentleman

**References**


**See Also**

`randomGraph`, `randomEGraph`
removeEdge

Examples

```r
set.seed(123)
c1 <- c(a = 1, b = 1, c = 2, d = 4)

(g1 <- randomNodeGraph(c1))
stopifnot(validObject(g1))
```

Description

A function to remove the specified edges from a graph.

Usage

```r
removeEdge(from, to, graph)
```

Arguments

- **from**: from edge labels
- **to**: to edge labels
- **graph**: a graph object

Details

A new graph instance is returned with the edges specified by corresponding elements of the `from` and `to` vectors removed. If `from` and `to` are not the same length, one of them should have length one. All edges to be removed must exist in `graph`.

Value

A new instance of a graph with the same class as `graph` is returned with the specified edges removed.

Author(s)

R. Gentleman

See Also

`addNode`, `addEdge`, `removeNode`
Examples

V <- LETTERS[1:4]
edL1 <- vector("list", length=4)
names(edL1) <- V
for(i in 1:4)
edL1[[i]] <- list(edges=c(2,1,4,3)[i], weights=sqrt(i))
gR <- graphNEL(nodes=V, edgeL=edL1)

gX <- removeEdge("A", "B", gR)

set.seed(123)
g <- randomEGraph(V=letters[1:5], edges=5)
g2 <- removeEdge(from=c("a","b"), to=c("c","e"), g)

Description

A function to remove a node from a graph. All edges to and from the node are also removed.

Usage

removeNode(node, object)

Arguments

node The label of the node to be removed.
object The graph to remove the node from.

Details

The specified node is removed from the graph as are all edges to and from that node. A new instance of the same class as object with the specified node(s) is returned.

Note, node can be a vector of labels, in which case all nodes are removed.

This is similar to subGraph.

Value

A new instance of a graph of the same class as object but with all specified nodes removed.

Author(s)

R. Gentleman

See Also

removeEdge, addEdge, addNode, subGraph
**Examples**

```r
V <- LETTERS[1:4]
edL2 <- vector("list", length=4)
names(edL2) <- V
for(i in 1:4)
edL2[[i]] <- list(edges=c(2,1,2,1)[i], weights=sqrt(i))
gR2 <- graphNEL(nodes=V, edgeL=edL2, edgemode="directed")
gX <- removeNode("C", gR2)
```

---

**renderInfo-class**  

Class "renderInfo"

**Description**

A container class to manage graph rendering attributes.

**Objects from the Class**

Objects can be created by calls of the form `new("renderInfo")` or by using the initializer `.renderInfoPrototype`.

**Slots**

- `pars`: List of default rendering attributes with two items `nodes` and `edges`. When not set further down the parameter hierarchy, these defaults will be used for all nodes/edges in the graph.
- `nodes`: Named list of attributes specific to nodes.
- `edges`: Named list of attributes specific to edges.
- `graph`: Named list of graph-wide attributes.

Each item of nodes and edges can take arbitrary vectors, the only restriction is that they have to be of either length 1 or length equal to the number of nodes or edges, respectively.

`pars` and `graph` can take arbitrary scalars, the latter for both edges and nodes.

**Methods**

The following are functions rather than methods and build the API to control the graphical output of a graph when it is plotted using `renderGraph`.

- `parRenderInfo`, `parRenderInfo<-`: getter and setter for items of slot `pars`
- `nodeRenderInfo`, `nodeRenderInfo<-`: getter and setter for items of slot `nodes`
- `edgeRenderInfo`, `edgeRenderInfo<-`: getter and setter for items of slot `edges`
- `graphRenderInfo`, `graphRenderInfo<-`: getter and setter for items of slot `graph`

The getters all take two arguments: `g` is a graph object and `name` is a character giving the name of one of the item in the respective slot. When `name` is missing this will give you the whole list.

The setters are a bit more complex: `nodeRenderInfo<-` and `edgeRenderInfo<-` can take
**named list of named vectors** where the names have to match the node or edge names. Items in the vector that don’t match a valid edge or node name will be silently ignored. For undirected edges the order of head nodes and tail nodes in edge names is ignored, i.e. `a~b` is equivalent to `codeb~a`.

**named list of skalars** which will set all the attribute for all edges or nodes in the graph `parRenderInfo<~` will only take a list with items `nodes`, `edges` and `graph`. The content of these list items can be arbitrary named vectors. `parRenderInfo<~` takes an arbitrary list.

Available rendering parameters for nodes are:

- **col**: the color of the line drawn as node border. Defaults to `black`.
- **lty**: the type of the line drawn as node border. Defaults to `solid`. Valid values are the same as for the R's base graphic parameter `lty`.
- **lwd**: the width of the line drawn as node border. Defaults to 1. Note that the underlying low level plotting functions do not support vectorized `lwd` values. Instead, only the first item of the vector will be used.
- **fill**: the color used to fill a node. Defaults to `transparent`.
- **textCol**: the font color used for the node labels. Defaults to `black`.
- **fontsize**: the font size for the node labels in points. Defaults to 14. Note that the fontsize will be automatically adjusted to make sure that all labels fit their respective nodes. You may want to increase the node size by supplying the appropriate layout parameters to `Graphviz` in order to allow for larger fontsizes.
- **cex**: Expansion factor to further control the fontsize. As default, this parameter is not set, in which case the fontsize will be clipped to the node size. This mainly exists to for consistency with the base graphic parameters and to override the clipping of fontsize to nodesize.

Available rendering parameters for edges are:

- **col**: the color of the edge line. Defaults to `black`.
- **lty**: the type of the edge line. Defaults to `solid`. Valid values are the same as for the R’s base graphic parameter `lty`.
- **lwd**: the width of the edge line. Defaults to 1.
- **textCol**: the font color used for the edge labels. Defaults to `black`.
- **fontsize**: the font size for the edge labels in points. Defaults to 14.
- **cex**: Expansion factor to further control the fontsize. This mainly exists to be consistent with the base graphic parameters.

**Author(s)**

Deepayan Sarkar, Florian Hahne
Examples

```r
g <- randomGraph(letters[1:4], 1:3, p=0.8)
nodeRenderInfo(g) <- list(fill=c("a"="red", "b"="green"))
edgeRenderInfo(g) <- list(lwd=3)
edgeRenderInfo(g) <- list(lty=3, col="red")
parRenderInfo(g) <- list(edges=list(lwd=2, lty="dashed"), nodes=list(col="gray", fill="gray"))
nodeRenderInfo(g)
edgeRenderInfo(g, "lwd")
edgeRenderInfo(g, c("lwd", "col"))
parRenderInfo(g)
```

reverseEdgeDirections  Reverse the edges of a directed graph

Description

Return a new directed graph instance with each edge oriented in the opposite direction relative to the corresponding edge in the input graph.

Usage

```r
reverseEdgeDirections(g)
```

Arguments

g A graph subclass that can be coerced to graphAM

Details

WARNING: this doesn’t handle edge attributes properly. It is a preliminary implementation and subject to change.

Value

A graphNEL instance

Author(s)

S. Falcon

Examples

```r
g <- graphNEL(nodes=c("a", "b", "c"),
edgeL=list(a=c("b", "c"), b=character(0), c=character(0)),
edgemode="directed")
stopifnot(isAdjacent(g, "a", "b"))
stopifnot(!isAdjacent(g, "b", "a"))
```
grev <- reverseEdgeDirections(g)
stopifnot(!isAdjacent(grev, "a", "b"))
stopifnot(isAdjacent(grev, "b", "a"))

dsimpleEdge-class  
Class "simpleEdge".

Description
A simple class for representing edges in graphs.

Objects from the Class
Objects can be created by calls of the form new("simpleEdge", ...).

Slots
edgeType: Object of class "character"; the type of the edge.
weight: Object of class "numeric"; the edge weight.
directed: Object of class "logical"; is the edge directed.
bNode: Object of class "character"; the beginning node.
eNode: Object of class "character"; the ending node.

Methods
No methods defined with class "simpleEdge" in the signature.

Note
All slots are length one vectors (this is not currently checked for). If the edge is not directed there is no real meaning to the concepts of beginning node or ending node and these should not be interpreted as such.

Author(s)
R. Gentleman

Examples
new("simpleEdge", bNode="A", eNode="D")
Standard labeling of edges with integers

**Description**

Functions to convert between from-to representation and standard labeling of the edges for undirected graphs with no self-loops.

**Usage**

```
ftM2int(ft)
int2ftM(i)
```

**Arguments**

- `i` Numeric vector.
- `ft` Numeric nx2 or 2xn matrix.

**Details**

A standard 1-based node labeling of a graph G=(V,E) is a one-to-one mapping between the integers from 1 to |V| and the nodes in V. A standard 1-based edge labeling of an undirected graph G=(V,E) with no self-loops is the one-to-one mapping between the integers from 1 to |V| choose 2 = |V|*(|V|-1)/2 such that the edge labeled 1 is between nodes 2 and 1, the edge labeled 2 is between nodes 3 and 1, the edge labeled 3 is between nodes 3 and 2, and so on.

**Value**

For `ftM2int`, a numeric vector of length n. For `int2ftM`, a length(i) x 2 matrix.

**Author(s)**

Wolfgang Huber

**Examples**

```r
nNodes <- 200
nEdges <- choose(nNodes, 2)
i <- 1:nEdges
ft <- int2ftM(i)
ft[1:6,]
stopifnot(all(ft[,1]>ft[,2])) ## always from higher to lower
stopifnot(!any(duplicated(paste(ft[,1], ft[,2]))))
stopifnot(ft[nEdges, 1]==nNodes, ft[nEdges, 2]==nNodes-1)

j <- ftM2int(ft)
stopifnot(all(i==j))
```
subGraph

Create a Subgraph

Description

Given a set of nodes and a graph this function creates and returns subgraph with only the supplied nodes and any edges between them.

Usage

subGraph(snodes, graph)

Arguments

- **snodes**: A character vector of node labels.
- **graph**: A graph object, it must inherit from the graph class.

Details

The returned subgraph is a copy of the graph. Implementations for Implementations for graphNEL, distGraph and clusterGraph.

Value

A graph of the same class as the graph argument but with only the supplied nodes.

Author(s)

R. Gentleman

See Also

nodes, edges

Examples

```r
set.seed(123)
x <- rnorm(26)
names(x) <- letters
library(stats)
d1 <- dist(x)
g1 <- new("distGraph", Dist=d1)
subGraph(letters[1:5], g1)
```
Description

There are two basic methods of generating dot (http://www.graphviz.org) language serializations of R graph-class structures. First, using the toDot methods of the Rgraphviz package, the native graphviz agraph-associated methods can be employed to create the dot serialization. Second, with the methods described here, R functions can be used to perform the serialization directly from the graph data structure, without Rgraphviz.

Methods

G = "graphNEL", outDotFile = "character", renderList = "list", optList = "list" create dot language description of graph

G = "graphNEL", outDotFile = "character", renderList = "missing", optList = "missing" create dot language description of graph

G = "graphNEL", outDotFile = "character", renderList = "missing", optList = "list" create dot language description of graph

G = "graphNEL", outDotFile = "missing", renderList = "missing", optList = "missing" create dot language description of graph

G = "graphNEL", outDotFile = "missing", renderList = "missing", optList = "list" create dot language description of graph

G = "graphNEL", outDotFile = "missing", renderList = "list", optList = "missing" create dot language description of graph

G = "graphNEL", outDotFile = "missing", renderList = "list", optList = "list" create dot language description of graph

G = "compoundGraph", outDotFile = "character", renderList = "list", optList = "missing" create dot language description of graph

G = "compoundGraph", outDotFile = "character", renderList = "list", optList = "list" create dot language description of graph

G = "compoundGraph", outDotFile = "missing", renderList = "list", optList = "missing" create dot language description of graph

See Also

toDot-methods
Examples

```r
eample(randomGraph)
tmp <- tempfile()
toDotR( g1, tmp )
readLines(tmp)
unlink(tmp)
```

toDotWithRI

*Translate a graph to "dot" including rendering information*

Description

The function takes a graph object and translates it into the dot format. All rendering information is written verbatim into the dot graph as well.

Usage

```r
toDotWithRI(graph, graph_name = NULL, subGraphList = list(), isStrict = TRUE)
```

Arguments

- `graph`: An object of graph
- `graph_name`: The name of the graph
- `subGraphList`: A list of objects of class graph. Each of these is used as a subgraph in the graph. The names in the list are taken as the names of the subgraph. If a graph is to be seen as a cluster (i.e. with a box around it), its name has to start with `cluster`
- `isStrict`: Should the graph be strict

Details

Given a graph object, it is translated into the dot language so that it can be rendered using the graphviz software. In addition to plotting the graph itself, all the rendering information is being used as well.

- `graphRenderInfo` attributes are written as an attribute list after the graph statement in dot.
- `nodeRenderInfo` attributes are written as attribute lists after each node. If an attribute is constant across all node, a global node attribute is written instead of many individual ones. Newlines in attributes do not lead to newlines in labels. In `label`, `headlabel` and `taillabel`, in order to get a newline, right justification or left justification, the two character sequences `\n`, `\r` and `\l` have to be written (i.e. in order to create this in R, the backslash has to be escaped in a string, that is has to be written as a double-backslash).
- `edgeRenderInfo` attributes as written as attribute lists after each edge, unless an attribute is constant, then it is written as a global edge attribute.

In general, all attribute values are being wrapped in double-quotes, unless the attribute value start with a `<` and ends with a `>`. In this case it is taken as html content and not wrapped in double quotes (nor are contained newlines escaped).

The resulting graph in dot format is returned as a character vector.
Value

A character vector with the graph in dot format

Author(s)

Holger Hoefling <hhoeflin@gmail.com>

Description

For a directed graph the underlying graph is the graph that is constructed where all edge orientation is ignored. This function carries out such a transformation on graphNEL instances.

Usage

ugraph(graph)

Arguments

graph a graph object.

Details

If \texttt{graph} is already undirected then it is simply returned.

If \texttt{graph} is a multi-graph (has multiple edges) an error is thrown as it is unclear how to compute the underlying graph in that context.

The method will work for any graph subclass for which an edgeMatrix method exists.

Value

An instance of graphNEL with the same nodes as the input but which is undirected.

Author(s)

R. Gentleman

References

Graph Theory and its Applications, J. Gross and J. Yellen.

See Also

connComp edgeMatrix
validGraph

Test whether graph object is valid

Description

validGraph is a validating function for a graph object.

Usage

validGraph(object, quietly=FALSE)

Arguments

object a graph object to be tested
quietly TRUE or FALSE indicating whether output should be printed.

Value

If the graph object is valid, TRUE is returned otherwise FALSE is returned. If object is not a valid graph and quietly is set to FALSE then descriptions of the problems are printed.

Author(s)

Elizabeth Whalen

See Also

graph-class

Examples

testGraph<-graphNEL()
testGraph@nodes<-c("node1","node2","node3")
validGraph(testGraph)
**write.tlp**

Write a graph object in a file in the Tulip format

---

**Description**

Write a graph object in a file in the Tulip format.

**Usage**

```python
write.tlp(graph, filename)
```

**Arguments**

- `graph`: a graph object
- `filename`: Name of the output file

**Details**

The Tulip format is used by the program Tulip.

**Author(s)**

Laurent Gautier <laurent@cbs.dtu.dk>

**References**

http://www.tulip-software.org/
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