# Package 'pathRender'

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Version 1.77.0 Title Render molecular pathways Author LiLong <lilong@isb-sib.ch> Maintainer Vince Carey <stvjc@channing.harvard.edu> Depends graph, Rgraphviz, RColorBrewer, cMAP, AnnotationDbi, methods, stats4 Suggests ALL, hgu95av2.db Description build graphs from pathway databases, render them by Rgraphviz. License LGPL URL http://www.bioconductor.org LazyLoad yes biocViews GraphAndNetwork, Pathways, Visualization git\_url https://git.bioconductor.org/packages/pathRender git\_branch devel git\_last\_commit 1287e36 git\_last\_commit\_date 2025-04-15 **Repository** Bioconductor 3.22

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coloredGraph-class Class "coloredGraph"

# Description

a graph to which color attributes have been attached

# **Objects from the Class**

Objects can be created by calls of the form new("coloredGraph", nodes, edgeL, edgemode). these are graphNEL instances with some additional graphData

# Slots

```
nodes: Object of class "vector" ~~
```

edgeL: Object of class "list" ~~

edgeData: Object of class "attrData" ~~

nodeData: Object of class "attrData" ~~

renderInfo: Object of class "renderInfo" ~~

graphData: Object of class "list" ~~

# Extends

Class "graphNEL-class", directly. Class "graph-class", by class "graphNEL", distance 2.

#### Methods

plot signature(x = "coloredGraph"): ...

# Examples

```
showClass("coloredGraph")
example(randomGraph)
nn = nodes(g1)
x = runif(length(nn))
names(x) = nn
h1 = colorNodes(g1, x, colorRampPalette(brewer.pal(9, "Blues"))(length(nn)),
pwayRendAttrs)
h1
plot(h1)
```

colorNodes

#### Description

attach node coloring information to a graphNEL instance

# Usage

colorNodes(g, nodeAss, pal, attgen)

# Arguments

g	graphNEL instance
nodeAss	color map for nodes: vector with elements evaluating to colors and nodes as element names
pal	a palette (use colorRampPalette for color interpolation)
attgen	attribute generating function – pwayRendAttrs is prototype

# Value

a graphNEL instance with additional rendering data

## Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# Examples

```
example(randomGraph)
nn = nodes(g1)
x = runif(length(nn))
names(x) = nn
h1 = colorNodes(g1, x, colorRampPalette(brewer.pal(9, "Blues"))(length(nn)),
pwayRendAttrs)
h1
```

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obtain a graph object corresponding to a cMAP pathway

# Description

obtain a graph object corresponding to a cMAP pathway

# Usage

graphcMAP(pname)

#### Arguments

pname

character token identifying a KEGG or cMAP pathway

# Details

reuses code from pathRender but emits a graphNEL-class instance with some additional information for rendering

## Value

an instance of pwayGraph, which extends graphNEL

# Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# Examples

```
G1 = graphcMAP("p53pathway")
G1
nodes(G1)
if (require(Rgraphviz)) plot(G1)
```

plotExGraph	plot a gene network, coloring nodes according to relative expression
	values

# Description

plot a gene network, coloring nodes according to relative expression values

# Usage

```
plotExGraph(g, es, sampind=1, pal=colorRampPalette(brewer.pal(9, "Blues"))(length(nodes(g))), attg
```

#### Arguments

g	graph representing a gene network
es	an ExpressionSet instance
sampind	sample to be used to obtain relative expression values
pal	palette for coloring the nodes
attgen	attribute generating function

# Details

plots a colored network on the current graphics display

# Value

as returned by Rgraphviz plot method for graphNEL instances

#### pwayGraph-class

#### Author(s)

Vince Carey <stvjc@channing.harvard.edu>

#### Examples

```
library(graph)
data(pancrCaIni)
library(ALL)
data(ALL)
library(hgu95av2.db)
collap1 = reduceES( ALL, nodes(pancrCaIni), revmap(hgu95av2SYMBOL), "symbol", mean )
library(RColorBrewer)
plotExGraph( pancrCaIni, collap1, 1 )
```

pwayGraph-class Class "pwayGraph" – extension to graphNEL for pathway rendering

# Description

extension to graphNEL for pathway rendering

### **Objects from the Class**

Objects can be created by calls of the form new("pwayGraph", nodes, edgeL, edgemode). There is a plot method that will work reasonably well if the plotting surface is big enough.

### Slots

pwaySource: Object of class "character" KEGG or BIOCARTA

nodes: Object of class "vector" pathway constituents in the native vocabulary

edgeL: Object of class "list" constituent relations in the native vocabulary

edgeData: Object of class "attrData" relationship attributes

nodeData: Object of class "attrData" node attributes

renderInfo: Object of class "renderInfo" render info

graphData: Object of class "list" this holds the special rendering attributes for edges and nodes, for nodes it seems particularly important to have fixedsize = FALSE

# Extends

Class graphNEL-class, directly. Class graph-class, by class "graphNEL", distance 2.

#### Methods

plot signature(x = "pwayGraph"): renders the pathway

# Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# Examples

```
showClass("pwayGraph")
G1 = graphcMAP("stresspathway")
G1@graphData$nAttrs$labels[1:10]
```

reduceES

collapse the assay values in an ExpressionSet to a set of specified genes, using a statistic when multiple probes map to a given gene

# Description

collapse the assay values in an ExpressionSet to a set of specified genes, using a statistic when multiple probes map to a given gene

# Usage

reduceES(es, annovec, ann2featMap, pdvname="symbol", collapseFun=NULL)

# Arguments

es	ExpressionSet instance
annovec	genes to retain
ann2featMap	either an AnnDbBimap from AnnotationDbi (typically constructed with revmap(), or a named vector mapping from symbols to probe set IDs
pdvname	featureData variable name to be used to hold the annotations of variables kept
collapseFun	statistical function for collapsing data across probes mapping to the same gene

# Value

An ExpressionSet instance limited to genes in annovec, condensed if necessary using collapseFun to get one number per gene from multiple probes

# Author(s)

Vince Carey <stvjc@channing.harvard.edu>

# Examples

```
library(ALL)
data(ALL)
library(hgu95av2.db)
rr = revmap(hgu95av2SYMBOL)
exprs(reduceES(ALL[,1:3], c("BCL2", "CPNE1"), rr, "sym", mean))
```

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rendercMAPPathway Render pathways from cMAP

# Description

Build graphs based on pathway or interaction data from cMAP database, render them using Rgraphviz.

# Usage

rendercMAPPathway(pname, ino=0)

#### Arguments

pname	name of the pathway to render
ino	index of the interaction in the given pathway to render

# Details

For a given pathway in cMAP database, we build a subgraph for each interaction in the pathway, join them together to form the graph for the complete pathway. The subgraphs for interactions and the graph for the pathway include info for rendering, such as labels/shapes/fillcolors for nodes, colors/styles/weights for edges. If user specifies an index of interaction, only the interaction is rendered. Otherwise, the complete pathway is rendered.

#### Value

None. A graphical output is presented.

#### Author(s)

Li Long <li.long@isb-sib.ch>

# Examples

```
rendercMAPPathway("plateletapppathway")
rendercMAPPathway("plateletapppathway", 5)
rendercMAPPathway("hsa00601")
rendercMAPPathway("hsa00601", 10)
```

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