# Package 'graphite'

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Title GRAPH Interaction from pathway Topological Environment

**Description** Graph objects from pathway topology derived from KEGG, Panther, PathBank, PharmGKB, Reactome SMPDB and WikiPathways databases.

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URL https://github.com/sales-lab/graphite

BugReports https://github.com/sales-lab/graphite/issues

**Depends** R (>= 4.2), methods

- **Imports** AnnotationDbi, graph (>= 1.67.1), httr, rappdirs, stats, utils, graphics, rlang, purrr
- Suggests checkmate, a4Preproc, ALL, BiocStyle, codetools, hgu133plus2.db, hgu95av2.db, impute, knitr, org.Hs.eg.db, parallel, R.rsp, RCy3, rmarkdown, SPIA (>= 2.2), testthat, topologyGSA (>= 1.4.0)
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VignetteBuilder R.rsp

- **biocViews** Pathways, ThirdPartyClient, GraphAndNetwork, Network, Reactome, KEGG, Metabolomics
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as.list.PathwayList Convertion of PathwayLists into lists.

# Description

Converts a PathwayList into a list of Pathways.

## Usage

## S3 method for class 'PathwayList'
as.list(x, ...)

# Arguments

| х | a PathwayList object       |
|---|----------------------------|
|   | extra arguments to as.list |

# Value

A list of pathways.

## Author(s)

Gabriele Sales

## See Also

PathwayList

## Examples

as.list(pathways("hsapiens", "kegg"))

buildPathway

#### Description

This function creates a new object of type Pathway given a data frame describing its edges.

## Usage

# Arguments

| id             | the pathway identifier.  |
|----------------|--|
| title          | the title of the pathway.  |
| species        | the species the pathway belongs to.  |
| database       | the name of the database the pathway derives from.                             |
| proteinEdges   | a data.frame of edges between proteins (or genes).                             |
|                | Must have the following columns: src_type, src, dest_type, dest, direction and |
|                | type.  |
|                | Direction must be one of the two strings: "directed" or "undirected".          |
| metaboliteEdge | S  |
|                | interactions between metabolites.  |
|                | Can be NULL. Otherwise, it must have the same structure as proteinEdges.       |
| mixedEdges     | interactions between metabolites and proteins.                                 |
|                | Can be NULL. Otherwise, it must have the same structure as proteinEdges.       |
| timestamp      | when the pathway was annotated, by default the time buildPathway is called.    |

#### Value

A new Pathway instance.

convertIdentifiers Convert the node identifiers of a pathway.

# Description

Converts the node identifiers of pathways.

If the option Ncpus is set to a value larger than 1 and the package parallel is installed, the conversion procedure will automatically use multiple cores.

#### Usage

```
convertIdentifiers(x, to)
```

### Arguments

| х  | can be a list of pathways or a single pathway  |
|----|--|
| to | a string describing the type of the identifier. Can assume the values "entrez", "symbol" or the name of one of the columns provided by an Annotation package (for example, "UNIPROT"). |

### Value

A Pathway object.

#### See Also

Pathway

```
r <- pathways("hsapiens", "reactome")
convertIdentifiers(r$`mTORC1-mediated signalling`, "symbol")</pre>
```

cytoscapePlot

# Description

Renders the topology of a pathway as a Cytoscape graph.

#### Usage

```
cytoscapePlot(pathway, ..., cy.ver = 3)
```

# Arguments

| pathway | a Pathway object.  |
|---------|--|
|         | optional arguments forwarded to pathwayGraph.                            |
| cy.ver  | select a Cytoscape version. Only version 3 is supported in this release. |

### Details

Requires the RCy3 package.

# Value

An invisible list with two items:

| graph | the graphNEL object sent to Cytoscape. |
|-------|--|
| suid  | the RCy3 network SUID.                 |

# See Also

Pathway

pathwayGraph

```
## Not run:
    r <- pathways()
    cytoscapePlot(convertIdentifiers(reactome$`Unwinding of DNA`, "symbol"))
## End(Not run)
```

Pathway-class

#### Description

A biological pathway.

#### Variants

A Pathway instance actually stores multiple variants of the same biological data.

This is the list of included variants:

- proteins: includes only interactions among proteins;
- metabolites: includes only interactions among metabolites;
- mixed: includes all available interactions.

#### Methods

pathwayId(p): Returns the native ID of the pathway.

pathwayTitle(p): Returns the title of the pathway.

pathwayDatabase(p): Returns the name of the database the pathway was derived from.

pathwaySpecies(p): Returns the name of the species in which the pathway was annotated.

pathwayTimestamp(p): Returns the date of pathway data retrieval.

pathwayURL(p): Returns the URL of the pathway in its original database, if available.

convertIdentifiers(p, to): Returns a new pathway using a different type of node identifiers.

The option which selects the desired pathway variant (see section "Variants" above).

If stringsAsFactors is TRUE, strings are converted to factors.

nodes(p, which = c("proteins", "metabolites", "mixed")): Returns the names of the nodes
 belonging to this pathway.

The option which selects the desired pathway variant (see section "Variants" above).

- plot(p): Shows the pathway topology in Cytoscape.
- runClipper(p, expr, classes, method, ...): Runs a clipper analysis over the pathway.
- runTopologyGSA(p, test, exp1, exp2, alpha, ...): Runs a topologyGSA analysis over the pathway.

#### Author(s)

Gabriele Sales

#### See Also

pathways

## pathwayDatabases

# Examples

```
reactome <- pathways("hsapiens", "reactome")
pathway <- reactome[[1]]
pathwayTitle(pathway)
pathwaySpecies(pathway)</pre>
```

nodes(pathway) edges(pathway)

pathwayDatabases List the available pathway databases.

# Description

Obtains the list of pathway databases available through graphite.

# Usage

```
pathwayDatabases()
```

# Value

Returns a data.frame with two columns: species and database.

## Author(s)

Gabriele Sales

# See Also

pathways

# Examples

pathwayDatabases()

pathwayGraph

# Description

Builds a graphNEL object representing the topology of a pathway.

## Usage

```
pathwayGraph(pathway, which = "proteins", edge.types = NULL)
```

# Arguments

| pathway    | a Pathway object.   |
|------------|---|
| which      | the pathway variant you want.                                   |
|            | See Pathway documentation for a list of the supported variants. |
| edge.types | keep only the edges maching the type names in this vector.      |

# Value

A graphNEL object.

#### See Also

#### Pathway

graphNEL

# Examples

```
r <- pathways("hsapiens", "reactome")
pathwayGraph(r$`mTORC1-mediated signalling`, edge.types="Binding")</pre>
```

PathwayList-class Class "PathwayList"

## Description

A collection of pathways from a single database.

# Extends

Class "Pathways", directly.

#### pathways

#### Methods

- 1[i] returns a selection of the pathways contained in the pathway list.
- 1[[i]] gives access to one of the pathways contained in the pathway list.
- 1\$'title' loads a pathways by its title.
- convertIdentifiers(1, to) returns a new list of pathways using a different type of node identifiers.
- length(1) returns the number of pathways contained in the list.
- names(1) returns the titles of the pathways contained in the list.
- prepareSPIA(1, pathwaySetName, print.names=FALSE) prepares the pathways for a SPIA analysis.
- runClipper(1, expr, classes, method, maxNodes=150, ...) runs a clipper analysis over all the pathways in the list.
- runTopologyGSA(1, test, exp1, exp2, alpha, maxNodes=150, ...) runs a topologyGSA analysis over all the pathways in the list.

## Author(s)

Gabriele Sales

#### See Also

pathways

pathways

Retrieve a list of pathways.

#### Description

Retrieve a list of pathways from a database for a given species. graphite currently supports the following databases:

- KEGG
- PANTHER
- PathBank
- PharmGKB
- Reactome
- SMPDB
- · WikiPathways

Call the pathwayDatabase function for more details.

#### Usage

pathways(species, database)

# Arguments

| species  | one of the supported species     |
|----------|----------------------------------|
| database | the name of the pathway database |

# Value

A PathwayList object.

# See Also

PathwayList, pathwayDatabases

# Examples

pathways("hsapiens", "reactome")

Pathways-class Class "Pathways"

## Description

A virtual class acting as a common parent to all other classes representing pathway databases.

# **Objects from the Class**

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

### Methods

No methods defined with class "Pathways" in the signature.

# Author(s)

Gabriele Sales

# See Also

PathwayList

prepareSPIA

#### Description

Prepare pathway dataset needed by runSPIA. See runSPIA and spia for more details.

#### Usage

```
prepareSPIA(db, pathwaySetName, print.names = FALSE)
```

#### Arguments

| db             | a PathwayList object or a list of Pathways.     |
|----------------|---|
| pathwaySetName | name of the output pathway set.                 |
| print.names    | print pathway names as the conversion advances. |

#### Value

This function has no return value.

#### References

Tarca AL, Draghici S, Khatri P, Hassan SS, Mittal P, Kim JS, Kim CJ, Kusanovic JP, Romero R. A novel signaling pathway impact analysis. Bioinformatics. 2009 Jan 1;25(1):75-82.

Adi L. Tarca, Sorin Draghici, Purvesh Khatri, et. al, A Signaling Pathway Impact Analysis for Microarray Experiments, 2008, Bioinformatics, 2009, 25(1):75-82.

Draghici, S., Khatri, P., Tarca, A.L., Amin, K., Done, A., Voichita, C., Georgescu, C., Romero, R.: A systems biology approach for pathway level analysis. Genome Research, 17, 2007.

### See Also

```
runSPIA
```

spia

PathwayList

runSPIA

#### Description

Run a topological analysis on an expression dataset using SPIA.

#### Usage

runSPIA(de, all, pathwaySetName, ...)

#### Arguments

| de             | A named vector containing log2 fold-changes of the differentially expressed genes. The names of this numeric vector are Entrez gene IDs.   |
|----------------|--|
| all            | A vector with the Entrez IDs in the reference set. If the data was obtained from<br>a microarray experiment, this set will contain all genes present on the specific<br>array used for the experiment. This vector should contain all names of the 'de'<br>argument. |
| pathwaySetName | The name of a pathway set created with prepareSPIA.  |
|                | Additional options to pass to spia.  |

#### Details

The spia option "organism" is internally used. It is an error use it in the additional options.

#### Value

The same of spia, without KEGG links. A data frame containing the ranked pathways and various statistics: pSize is the number of genes on the pathway; NDE is the number of DE genes per pathway; tA is the observed total preturbation accumulation in the pathway; pNDE is the probability to observe at least NDE genes on the pathway using a hypergeometric model; pPERT is the probability to observe a total accumulation more extreme than tA only by chance; pG is the p-value obtained by combining pNDE and pPERT; pGFdr and pGFWER are the False Discovery Rate and respectively Bonferroni adjusted global p-values; and the Status gives the direction in which the pathway is perturbed (activated or inhibited).

## References

Tarca AL, Draghici S, Khatri P, Hassan SS, Mittal P, Kim JS, Kim CJ, Kusanovic JP, Romero R. A novel signaling pathway impact analysis. Bioinformatics. 2009 Jan 1;25(1):75-82.

Adi L. Tarca, Sorin Draghici, Purvesh Khatri, et. al, A Signaling Pathway Impact Analysis for Microarray Experiments, 2008, Bioinformatics, 2009, 25(1):75-82.

Draghici, S., Khatri, P., Tarca, A.L., Amin, K., Done, A., Voichita, C., Georgescu, C., Romero, R.: A systems biology approach for pathway level analysis. Genome Research, 17, 2007.

## runTopologyGSA

#### See Also

spia

#### Examples

```
if (require(SPIA) && require(hgu133plus2.db)) {
   data(colorectalcancer)

   top$ENTREZ <- mapIds(hgu133plus2.db, top$ID, "ENTREZID", "PROBEID", multiVals = "first")
   top <- top[!is.na(top$ENTREZ) & !duplicated(top$ENTREZ), ]
   top$ENTREZ <- paste("ENTREZID", top$ENTREZ, sep = ":")
   tg1 <- top[top$adj.P.Val < 0.05, ]

   DE_Colorectal = tg1$logFC
   names(DE_Colorectal) <- tg1$ENTREZ
   ALL_Colorectal <- top$ENTREZ
   kegg <- pathways("hsapiens", "kegg")[1:20]
   kegg <- convertIdentifiers(kegg, "ENTREZID")
   prepareSPIA(kegg, "keggEx")
   runSPIA(de = DE_Colorectal, all = ALL_Colorectal, "keggEx")
   unlink("keggExSPIA.RData")
}</pre>
```

runTopologyGSA Run a topological analysis on an expression dataset using topologyGSA.

#### Description

Use graphical models to test the pathway components highlighting those involved in its deregulation.

If the option Ncpus is set to a value larger than 1 and the package parallel is installed, the conversion procedure will automatically use multiple cores.

# Usage

```
runTopologyGSA(x, test, exp1, exp2, alpha, ...)
```

#### Arguments

| x     | a PathwayList, a list of Pathways or a single Pathway object.            |
|-------|--|
| test  | Either "var" and "mean". Determine the type of test used by topologyGSA. |
| exp1  | Experiment matrix of the first class, genes in columns.                  |
| exp2  | Experiment matrix of the second class, genes in columns.                 |
| alpha | Significance level of the test.  |

... Additional parameters forwarded to topologyGSA. When invoked on a PathwayList, can use the named option "maxNodes" to limit the analysis to those pathways having up to this given number of nodes.

# Details

This function produces a warning and returns NULL when the number of genes in common between the expression matrices and the pathway is less than 3.

# Value

See documentation of pathway.var.test and pathway.mean.test.

#### References

Massa MS, Chiogna M, Romualdi C. Gene set analysis exploiting the topology of a pathway. BMC System Biol. 2010 Sep 1;4:121.

```
if (require(topologyGSA)) {
   data(examples)
   colnames(y1) <- paste("SYMBOL", colnames(y1), sep = ":")
   colnames(y2) <- paste("SYMBOL", colnames(y2), sep = ":")
   k <- pathways("hsapiens", "kegg")
   p <- convertIdentifiers(k[["Fc epsilon RI signaling pathway"]], "SYMBOL")
   runTopologyGSA(p, "var", y1, y2, 0.05)
}</pre>
```

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