

Package ‘categoryCompare’

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Version 1.55.0

Title Meta-analysis of high-throughput experiments using feature annotations

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URL <https://github.com/rmflight/categoryCompare>

BugReports <https://github.com/rmflight/categoryCompare/issues>

License GPL-2

Depends R (>= 2.10), Biobase, BiocGenerics (>= 0.13.8),

Suggests knitr, GO.db, KEGGREST, estrogen, org.Hs.eg.db, hgu95av2.db, limma, affy, genefilter, rmarkdown

Imports AnnotationDbi, hwriter, GSEABase, Category (>= 2.33.1), GOstats, annotate, colorspace, graph, RCy3 (>= 1.99.29), methods, grDevices, utils

LazyLoad yes

Description Calculates significant annotations (categories) in each of two (or more) feature (i.e. gene) lists, determines the overlap between the annotations, and returns graphical and tabular data about the significant annotations and which combinations of feature lists the annotations were found to be significant. Interactive exploration is facilitated through the use of RCytoscape (heavily suggested).

SystemRequirements Cytoscape (>= 3.6.1) (if used for visualization of results, heavily suggested)

TODO Text and HTML output without graphs.

biocViews Annotation, GO, MultipleComparison, Pathways, GeneExpression

VignetteBuilder knitr

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Contents

categoryCompare-package	3
breakEdges-methods	4
ccCompare-methods	5
ccCompareCollection-class	7
ccCompareGeneric-methods	8
ccCompareResult-class	8
ccData	9
ccEnrich-method	10
ccEnrichCollection-class	12
ccEnrichResult-class	13
ccGeneList-class	14
ccOptions-class	16
ccOutCyt-methods	17
ccSigList-class	18
cwReload-methods	19
cytOutData-methods	20
cytOutNodes-methods	21
fdr	21
GENccEnrichResult-class	22
getGeneSymbol	23
graphType-methods	24
HyperGParamsCC-class	25
HyperGResultCC-class	26
hyperGTestCC	27
listNames	28
mergedData-class	29
mergeLists-methods	30
minCount	31
minNodes	32
pvalueType	33
resetColors-methods	34
show-methods	34

Index

35

categoryCompare-package

Meta-analysis of high-throughput experiments using feature annotations

Description

Calculates significant annotations (categories) in each of two (or more) feature (i.e. gene) lists, determines the overlap between the annotations, and returns graphical and tabular data about the significant annotations and which combinations of feature lists the annotations were found to be significant. Interactive exploration is facilitated through the use of RCytoscape (heavily suggested).

Details

Package:	categoryCompare
Version:	1.21.2
License:	GPL-2
Depends:	Biobase (>= 1.15.29), AnnotationDbi (>= 0.1.15), Category
Suggests:	methods, GSEABase, hwriter, colorspace, graph, GO.db, KEGG.db, estrogen, org.Hs.eg.db, hgu95av
Imports:	Biobase (>= 1.15.29), AnnotationDbi (>= 0.1.15), hwriter, GSEABase, Category (>= 2.21.2), GStat
LazyLoad:	yes
biocViews:	Bioinformatics, Annotation, GO, MultipleComparisons, Pathways, GeneExpression
SystemRequirements:	Cytoscape (>= 2.8.0) (if used for visualization of results, heavily suggested), CytoscapeRPC plugin (
TODO:	Text and HTML output without graphs.
Built:	R 2.15.0; ; 2012-03-15 18:42:40 UTC; windows

Index:

GENccEnrichResult-class

Class '"GENccEnrichResult"'

HyperGParamsCC-class

Class "HyperGParamsCC"

HyperGResultCC-class

Class "HyperGResultCC"

breakEdges

Break Cytoscape (or graphNEL) Network Edges

breakEdges-methods

Methods for Function 'breakEdges' in Package
'categoryCompare'

categoryCompare-package

Meta-analysis of high-throughput experiments
using feature annotations

ccCompare-methods

Comparison of enriched annotations

ccCompareCollection-class

Class '"ccCompareCollection"'

ccCompareGeneric-methods

Methods for Function 'ccCompareGeneric' in
Package 'categoryCompare'

ccCompareResult-class

Class '"ccCompareResult"'

ccData	Test data for 'categoryCompare'
ccEnrich-method	Perform annotation enrichment for multiple gene lists
ccEnrichCollection-class	Class "ccEnrichCollection"
ccEnrichResult-class	Class "ccEnrichResult"
ccGeneList-class	Class "ccGeneList"
ccOptions-class	Class "ccOptions"
ccOutCyt-methods	Methods for Function 'ccOutCyt' in Package 'categoryCompare'
ccSigList-class	Class "'ccSigList'"
cwReload-methods	Methods for Function 'cwReload' in Package 'categoryCompare'
cytOutData-methods	Methods for Function 'cytOutData'
cytOutNodes-methods	Methods for Function 'cytOutNodes'
fdr	Number of FDR runs to perform
getGeneSymbol	Entrez to name, symbol, GO and path conversion, as well as general ID to ID conversion.
graphType-methods	graphType
hyperGTestCC	Hypergeometric testing with false discovery rate
listNames	listNames
mergeLists-methods	Function 'mergeLists' in Package 'categoryCompare'
mergedData-class	Class "'mergedData'"
minCount	minCount
minNodes	Delete nodes with less than a certain number of genes annotated
pvalueType	Type of p-values to return from object
resetColors-methods	resetColors
show-methods	Methods for Function show in Package 'categoryCompare'

Further information is available in the following vignettes:

categoryCompare_vignette categoryCompare: High-throughput data meta-analysis using gene annotations (source)

Author(s)

Robert M. Flight <rflight79@gmail.com>

Maintainer: Robert M. Flight <rflight79@gmail.com>

Description

Methods for function `breakEdges` in package **categoryCompare**

Methods

`signature(cwObject = "ccCompareResult", cutoff = "numeric")` Allows one to remove edges in the `ccCompareResult` `mainGraph` slot prior to passing it into Cytoscape for visualization. Given that the number of edges can be rather large (especially for Gene Ontology) this can easily speed up the transfer, without actually losing any information.

`signature(cwObject = "numeric", cutoff = "numeric")` Once an annotation graph is in Cytoscape, remove edges above or below the cutoff. Note that this does not affect the original graph in the `ccCompareResult` object.

Author(s)

Robert M Flight

See Also

[breakEdges ccCompareResult ccOutCyt](#)

Examples

```
data(ccData)

# breaking the edges in a ccCompareResult
ccResults$BP <- breakEdges(ccResults$BP, 0.8)
## Not run:
hasCy <- (if (.Platform$OS.type %in% "windows") {
  (length(grep("Cytoscape", system("tasklist", intern=TRUE))) > 0)})

if hasCy {

  cwObj <- ccOutCyt(ccResults$BP, ccOpts)
  # now breaking them in the CytoscapeWindow object
  breakEdges(cwObj, 0.85)
  Sys.sleep(10)
  RCy3::deleteWindow(cwObj)
}

## End(Not run)
```

Description

Takes the results from [ccEnrich](#) and compares the enriched annotations based on the settings previously set in [ccOptions](#). Returns a `ccCompareResult` or `ccCompareCollection` object, see Details.

Usage

```
ccCompare(ccEnrichResult, ccOptions)
```

Arguments

ccEnrichResult The enriched annotations collection returned from `ccEnrich`. This can be the `ccEnrichCollection`, `GOccEnrichResult`, or `KEGGccEnrichResult`

ccOptions A `ccOptions` object that will determine which lists to actually compare against each other. See details below.

Details

Based on the enrichments found for each gene list, we now want to compare the annotations between lists. `ccCompare` accesses the annotations for each enrichment performed for each list, and makes the comparisons defined in `ccOptions`.

Value

`ccCompare` generates both a graph of the comparisons (to show how the categories are linked to each list and each other) and tabular output. The tabular output is a data frame, with ID for each term that was considered as a candidate annotation for each list, as well as a long description (Desc) of what the term is, and then membership and statistics from each gene list.

For each type of comparison (GO, KEGG, etc) a `ccCompareResult` is generated, with the following slots:

<code>mainGraph</code>	Annotations arranged as a graph
<code>mainTable</code>	The tabular results from all enrichment calculations combined into one
<code>allAnnotation</code>	A list of lists, where each entry is the annotation identifier, then a list for each comparison, with the genes that are annotated to that term that also belong to each list

The default is to generate an overlap graph for GO and KEGG, where the overlap is a measure of the similarity of the features (genes) annotated to each annotation term (based on a formula from `EnrichmentMap`). Optionally for GO, one can generate a hierarchical layout where the parent GO terms of the significant terms will also be included in the graph, with term origin saved in the node annotation (see example below to do this).

Only those terms with more than 10 and less than 500 annotated genes (according to the GO annotation file) are included.

When using weighted overlap graphs and **RCy3** for viewing, it is recommended to use `breakEdges` and `minNodes` to remove edges with low weights and nodes with only a few genes from the dataset annotated to them.

Author(s)

Robert M Flight

See Also

[ccCompareResult](#) [ccCompareCollection](#) [ccOutCyt](#) [breakEdges](#) [outType](#) [ccEnrich](#)

Examples

```
## Not run:
require(GO.db)
require(KEGG.db)
require(org.Hs.eg.db)

## End(Not run)
data(ccData)

# note that enrichLists is generated from ccEnrich
# ccResults <- ccCompare(enrichLists,ccOpts)
ccResults

# use the GO hierarchy tree
graphType(enrichLists$BP) <- "hierarchical"
# ccResultsBPHier <- ccCompare(enrichLists$BP,ccOpts)
ccResultsBPHier
```

```
ccCompareCollection-class
      Class "ccCompareCollection"
```

Description

Holds multiple [ccCompareResult](#) objects.

Objects from the Class

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

These are not normally created by the user, but rather by [ccCompare](#) while performing the categorical comparisons for each type of category

Slots

```
.Data: Object of class "list"
names: Object of class "character"
```

Extends

Class "[namedList](#)", directly. Class "[list](#)", by class "[namedList](#)", distance 2. Class "[vector](#)", by class "[namedList](#)", distance 3. Class "[AssayData](#)", by class "[namedList](#)", distance 3.

Methods

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

Author(s)

Robert M Flight

See Also

[ccCompareResult ccCompare](#)

Examples

```
showClass("ccCompareCollection")
```

ccCompareGeneric-methods

*Methods for Function ccCompareGeneric in Package **categoryCompare***

Description

Methods for function ccCompareGeneric in package **categoryCompare**

Methods

```
signature(gccResult = "GENccEnrichResult", ccOptions = "ccOptions")
```

ccCompareResult-class *Class "ccCompareResult"*

Description

Holds the results from a single category comparison

Objects from the Class

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

Slots

mainGraph: Object of class "graph". Holds the graph describing the relationships between the annotations

subGraph: Object of class "list". Not currently used

mainTable: Object of class "data.frame". Table of results, with all the various statistics for each annotation in the category

allAnnotation: Object of class "list". For each annotation, which genes from which comparison are annotated to that particular annotation

categoryName: Object of class "character". Which category (e.g. GO, KEGG, etc) was used

ontology: Object of class "character". If GO, which ontology was used

Methods

allAnnotation signature(object = "ccCompareResult"): ...

mainGraph signature(object = "ccCompareResult"): ...

mainTable<- signature(object = "ccCompareResult"): ...

Author(s)

Robert M Flight

See Also

[ccCompare](#) [ccCompareCollection](#)

Examples

```
showClass("ccCompareResult")
```

ccData

Test data for categoryCompare

Description

Processed data from the estrogen example data set

Usage

```
data(ccData)
```

Format

table10: Log-ratio output from **limma** for the comparison of presence-absence of estrogen at 10 hours

table48: Log-ratio output from **limma** for the comparison of presence-absence of estrogen at 48 hours

gUniverse: All of the genes measured on the chip

gseaRes: Toy results of GSEA analysis of 3 different tissues

enrichLists: Apply [ccEnrich](#) to a ccGeneList from table10 and table48

ccResults: Apply [ccCompare](#) to enrichLists

ccResultsBPHier: Modify enrichLists\$BP to use a "hierarchical" layout

geneLists: a ccGeneList generated from genes in table10 and table48

ccOpts: a ccOptions object describing what we are going to do as far as feature list comparisons

Author(s)

Robert M Flight

Source

Taken from the **estrogen** package in Bioconductor, and then processed using the normal **affy** and **limma** tools.

See Also

[ccGeneList](#) [ccEnrichCollection](#) [ccCompareCollection](#) [ccEnrich](#) [ccCompare](#)

Examples

```
data(ccData)
```

ccEnrich-method

Perform annotation enrichment for multiple gene lists

Description

Takes a [ccGeneList](#) object containing all the information needed to perform enrichment calculations for Gene Ontology.

Usage

```
ccEnrich(ccGeneList)
```

Arguments

ccGeneList A [ccGeneList](#) object, which is really just a list of lists, with some extra slots to tell us how to examine results. Each entry in the list should be named to allow identification later on. Each sub list should contain a vector **genes** denoting the genes of interest, a vector **universe** denoting the gene background (i.e. all genes on the chip), and an entry **annotation** denoting an organism database package (such as **org.Hs.eg.db**). See [ccGeneList](#) for more details regarding this object.

Details

This function is essentially a wrapper for [hyperGTestCC](#) that performs all of the calculations for the many gene lists in one go, returning a list of [HyperGResultCC](#) objects, one for each of the **ccTypes** and each gene list. These various [HyperGResultCC](#) objects can then be accessed and results compared among the lists for each of the ontologies

Value

A list of [HyperGResultCC](#) objects, one for each **ccType** and gene list, returned as [ccEnrichResult](#) objects for each **ccType**. This can be passed with a [ccOptions](#) object to [ccCompare](#) to generate actual annotation comparisons.

Author(s)

Robert M Flight

See Also

[ccGeneList](#), [hyperGTestCC](#), [ccEnrichResult](#)

Examples

```
## Not run:
require(GO.db)
require(KEGG.db)
require(org.Hs.eg.db)

## End(Not run)
data(ccData)

g10 <- (unique(table10$Entrez[1:100]))
g48 <- (unique(table48$Entrez[1:100]))

list10 <- list(genes=g10, universe=gUniverse, annotation="org.Hs.eg.db")
list48 <- list(genes=g48, universe=gUniverse, annotation="org.Hs.eg.db")

geneLists <- list(T10=list10, T48=list48)
geneLists <- new("ccGeneList", geneLists, ccType=c("BP","KEGG"))
geneLists <- new("ccGeneList", geneLists, ccType=c("CC","KEGG"))

# set number of fdr runs to 0 to speed up runtime, not generally recommended.
```

```
geneLists <- new("ccGeneList", geneLists, ccType = c('BP', 'KEGG'), pvalueCutoff=0.01, fdr=0)
# enrichLists <- ccEnrich(geneLists)
```

ccEnrichCollection-class

Class "ccEnrichCollection"

Description

Holds multiple classes of [ccEnrichResult](#) in one object to allow [ccCompare](#) to work on only the one object and generate all of the results of a comparison.

Objects from the Class

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

Slots

.Data: Object of class "list"

names: Object of class "character" The names (generally GO ontologies or KEGG, but can be changed) of each set of results

Extends

Class "[namedList](#)", directly. Class "[list](#)", by class "namedList", distance 2. Class "[vector](#)", by class "namedList", distance 3. Class "[AssayData](#)", by class "namedList", distance 3.

Methods

pvalueCutoff<- signature(r = "ccEnrichCollection"): Changes the pvalueCutoff to be used to decide significant annotations for all of the contained ccEnrichResult objects

pvalueType<- signature(object = "ccEnrichCollection"): Changes whether to use p-values or fdr values to determine those annotations that are significant in all of the contained ccEnrichResult objects

minCount<- signature(object = "ccEnrichCollection"): how many features have to be annotated to a term to be reported as significant

graphType signature(object = "ccEnrichCollection"): Gets the type of graph that should be output for this collection

Author(s)

Robert M Flight

See Also

[ccEnrich](#) [hyperGTestCC](#) [ccCompare](#) [ccEnrichResult](#)

Examples

```
data(ccData)
enrichLists
```

ccEnrichResult-class *Class "ccEnrichResult"*

Description

Acts as a container object for multiple [HyperGResultCC](#) objects.

Objects from the Class

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

Extends

Class "[namedList](#)", directly. Class "[list](#)", by class "namedList", distance 2. Class "[vector](#)", by class "namedList", distance 3. Class "[AssayData](#)", by class "namedList", distance 3.

Methods

fdr signature(object = "ccEnrichResult"): get the number of runs using random feature lists were performed

pvalueCutoff signature(r = "ccEnrichResult"): what is the pvalueCutoff to determine significant annotations

pvalueCutoff<- signature(r = "ccEnrichResult"): change the pvalueCutoff for an annotation to be considered significant

pvalueType<- signature(object = "ccEnrichResult"): change whether p-values used are from "FDR" or raw p-values

minCount signature(object = "ccEnrichResult"): how many features need to belong to an annotation to be reported

minCount<- signature(object = "ccEnrichResult"): adjust the minCount

graphType signature(object = "ccEnrichResult"): what type of graph should be generated (generally set by the class of object)

graphType<- signature(object = "ccEnrichResult"): change the type of graph to generate by ccCompare

Author(s)

Robert M Flight

Examples

```
data(ccData)
enrichRes <- enrichLists[[1]]
fdr(enrichRes)
pvalueType(enrichRes)
enrichRes
pvalueType(enrichRes) <- 'pval'
enrichRes

pvalueCutoff(enrichRes)
pvalueCutoff(enrichRes) <- 0.01
enrichRes
```

ccGeneList-class

Class "ccGeneList"

Description

This stores the actual gene lists and related information that will be used in categoryCompare.

Objects from the Class

Objects can be created by calls of the form `new("ccGeneList", list)`. `ccGeneList` is actually just an extension of R list objects. The input list should be a list of lists. See Details for more information.

Slots

fdr: Object of class "numeric" The number of fdr runs to perform to account for different list sizes and term dependence

pvalueCutoff: Object of class "numeric" Value used to determine whether or not a particular term is significant or not

ccType: Object of class "character" What types of annotations to use. Currently supported ones include "BP", "MF", "CC" (from Gene Ontology) and "KEGG"

testDirection: Object of class "character" Are you interested in "over" or "under" represented annotations

Methods

fdr signature(object = "ccGeneList"): how many random runs to perform

fdr<- signature(object = "ccGeneList"): change the number of random runs

pvalueCutoff signature(object = "ccGeneList"): what is the pvalue to consider significant

pvalueCutoff<- signature(object = "ccGeneList"): change the cutoff for significance

ccType signature(object = "ccGeneList"): what type of annotations are going to be examined

ccType<- signature(object = "ccGeneList"): change the type of annotations to examine

testDirection signature(object = "ccGeneList"): query for "over" or "under" represented annotations

testDirection<- signature(object = "ccGeneList"): change the type of representation ("over" or "under")

listNames signature(object = "ccGeneList"): what are the names of the lists contained

Details

The input list should be a list of lists, with at least three sub-lists.

```
testList <- list(list1=list(genes='...',universe='...',annotation='...'), list2=list(...))
```

genes : These are the gene identifiers of the genes that are of interest (differentially expressed genes)

universe : All of the genes that were measured in this particular experiments (i.e. all the genes on the chip)

annotation : What organism or chip do these ID's come from (e.g. "org.Hs.eg.db" for Human Entrez gene ID's, "hgu133a.db" for probe ID's from the Affymetrix U133A chip)

data : A data-frame that contains extra information about the genes of interest. At the very least, the data-frame must have a column ID that matches the ID's contained in genes

What actually happens when running ccEnrich is that the appropriate HyperGParamsCC objects are generated for each geneList and each type of annotation (e.g. BP, CC, KEGG), and then the calculations performed on each one.

Note

The ccGeneList object is what will undergo all of the enrichment calculations. When the results are combined with the [ccOptions](#) object, we can get our results of actual comparisons between experiments.

Author(s)

Robert M Flight

See Also

[ccOptions](#)

Examples

```
data(ccData)
g10 <- (unique(table10$Entrez[1:100]))
g48 <- (unique(table48$Entrez[1:100]))

list10 <- list(genes=g10, universe=gUniverse, annotation="org.Hs.eg.db")
list48 <- list(genes=g48, universe=gUniverse, annotation="org.Hs.eg.db")

geneLists <- list(T10=list10, T48=list48)
geneLists <- new("ccGeneList", geneLists, ccType=c("BP","KEGG"))
geneLists
```

ccOptions-class	Class "ccOptions"
-----------------	-------------------

Description

These objects store the various options required by categoryCompare for actually making comparisons and generating output.

Objects from the Class

Objects can be created by calls of the form `new("ccOptions", listNames=c('list1', 'list2', etc))`. This is the minimum call required, and will generate a ccOptions object where comparisons are assumed between all the lists supplied. See the examples section for more examples of how to initialize new objects.

Slots

listNames: Object of class "character" The actual names of the various datasets defined in the ccData object

compareNames: Object of class "character" Which lists to compare, each entry should be a comma separated list

compareIndx: Object of class "list" List indices for each of the comparison, not usually set by the user. Generated automatically.

compareColors: Object of class "character" For graphical and tabular output each comparison can be colored. Should be one color for each comparison. Can be either an n by 3 matrix of rgb triples, or a character vector of hexadecimal color codes, or character vector of color names ('red', 'green', 'blue', etc)

cssClass: Object of class "character" Classnames used when generating HTML tables to color entries. Generated automatically upon initialization, or modifying compareNames

outType: Object of class "character" Sets the type of output generated by ccTables. Valid types are "html", "text", "rcy3" or "none", default is "text" when the ccOptions object is initialized without an outType specified.

Methods

compareColors signature(object = "ccOptions"): ...

compareColors<- signature(object = "ccOptions"): ...

compareIndx signature(object = "ccOptions"): ...

compareNames signature(object = "ccOptions"): ...

compareNames<- signature(object = "ccOptions"): ...

cssClass signature(object = "ccOptions"): ...

listNames signature(object = "ccOptions"): ...

listNames<- signature(object = "ccOptions"): ...

outType signature(object = "ccOptions"): ...

outType<- signature(object = "ccOptions"): ...

Author(s)

Robert M Flight

Examples

```

showClass("ccOptions")
## A very basic "ccOptions" for a comparison of two sets of data, "list1" and "list2"
c1 <- new("ccOptions", listNames=c('list1','list2'))
c1

## Now lets get a little more complicated
c1 <- new("ccOptions", listNames=c('list1','list2'),
compareNames=c('list1,list2','list1,list3'), compareColors=c('red','blue'))
c1

# set the type of output you want to eventually produce
c1 <- new("ccOptions", listNames=c('list1','list2'), outType='html')
c1

c1 <- new("ccOptions", listNames=c('list1','list2'), outType=c('html','text','none'))
c1

## Using RGB colors
ccCols <- matrix(c(255,0,0, 0,0,255), nrow=2, ncol=3)
ccCols <- rgb(ccCols, maxColorValue=255)
c1 <- new("ccOptions", listNames=c('list1','list2','list3'),
compareNames=c('list1,list2','list1,list3'), compareColors=ccCols)

## Using Hex colors
c1 <- new("ccOptions", listNames=c('list1','list2','list3'),
compareNames=c('list1,list2','list1,list3'), compareColors=c('#FF0000','#0000FF'))
c1

## or even using a color palette from R.
## Note that you need at least enough colors to cover all of individual and
## possible permutations (n!) if you use compareNames='all'
c1 <- new("ccOptions", listNames=c('list1','list2','list3'),
compareNames=c('list1,list2','list1,list3'), compareColors=rainbow(4))
c1

```

Description

Passes a `ccCompareResult` object to Cytoscape for interactive visualization of `ccCompare` results.

Details

Note that only some basic, required methods have been imported from RCy3 for use with categoryCompare, and these are hidden in the functions within categoryCompare and are not visible to the user. If access to all the functionality of RCytoscape is desired (and trust me, there is a lot of useful stuff in there), then the user should use `library(RCy3)` directly.

It should also be noted that deletion of edges via RCy3 is slow, so some edge filtering should be done by `breakEdges` prior to using `ccOutCyt`.

Methods

`signature(ccCompRes = "ccCompareResult", ccOpts = "ccOptions", ...)` At a minimum, this method requires a `ccCompareResult` and a `ccOptions` to work.

... may include:

layout = "character" to override the default layout set by `ccCompare`, as well as options

postText = "character" to add a user set string to the Cytoscape window

In addition, any of the arguments to `CytoscapeWindow` may also be set, such as host or port.

See Also

`ccCompareResult ccOptions ccCompare CytoscapeWindowClass`

Examples

```
## Not run:
hasCy <- (if (.Platform$OS.type %in% "windows") {
  (length(grep("Cytoscape", system("tasklist", intern=TRUE))) > 0)})

if hasCy {

  ccResults$BP <- breakEdges(ccResults$BP, 0.8)
  cwObj <- ccOutCyt(ccResults$BP, ccOpts)
  Sys.sleep(10)
  RCy3::deleteWindow(cwObj)
}

## End(Not run)
```

ccSigList-class

Class "ccSigList"

Description

Holds a generic list of significant annotations. Allows one to use Bioconductor annotation packages, or when combined into a `GENccEnrichResult`, use custom annotation / gene mappings.

Objects from the Class

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

Slots

sigID: Object of class "character"
categoryName: Object of class "character"
ontology: Object of class "character"
annotation: Object of class "character"

Methods

annotation signature(object = "ccSigList"): ...
category signature(object = "ccSigList"): ...
ontology signature(object = "ccSigList"): ...
sigID signature(object = "ccSigList"): ...

Author(s)

Robert M Flight

See Also

[GENccEnrichResult ccCompareGeneric](#)

Examples

```
showClass("ccSigList")
```

cwReload-methods

*Methods for Function cwReload in Package **categoryCompare***

Description

Methods for function cwReload in package **categoryCompare**

Methods

signature(oldCW = "numeric", windowName = "character", ccOpts = "ccOptions") This method is now deprecated as the cwObj doesn't store anything, but is merely the network SUID pointing to the network in Cytoscape. See RCy3::getNetworkSuid.

Description

Takes the saveObj generated by cytOutNodes and writes the data to a file

Value

A text file with the annotations previously saved using cytOutNodes

Methods

```
signature(saveObj = "list", compareResult = "ccCompareResult", mergedData = "mergedData")
  saveObj is the list object generated by cytOutNodes, compareResult is the object from
  ccCompare, and mergedData is created using mergeLists, but is optional.

... : optional arguments also include: orgType, default is "header" where each group is seperate,
      "annotate" pushes all the data into one table with a new column that designates which
      groups the annotation was found in; fileName, the name of a text file to output the results to;
      displayFile, whether or not to display the file (default is "FALSE")
```

Examples

```
## Not run:
hasCy <- (if (.Platform$OS.type %in% "windows") {
  (length(grep("Cytoscape", system("tasklist", intern=TRUE))) > 0)})

if hasCy {
  data(ccData)
  ccResults$BP <- breakEdges(ccResults$BP, 0.8)
  cwObj <- ccOutCyt(ccResults$BP, ccOpts)
  # user selects some nodes in Cytoscape
  RCy3::selectNodes(cwObj, c("GO:0007017", "GO:0000226", "GO:0007051", "GO:0007052"))
  savedNodes <- cytOutNodes("random1", cwObj) # save them
  # and selects some other nodes
  RCy3::selectNodes(cwObj,
    c("GO:0071103", "GO:0034728", "GO:0006323", "GO:0030261", "GO:0006334"),
    preserve.current.selection=FALSE)
  savedNodes <- cytOutNodes("random2", cwObj, savedNodes)

  # now spit results out to a file
  cytOutData(savedNodes, ccResults$BP)
}
## End(Not run)
```

cytOutNodes-methods *Methods for Function cytOutNodes*

Description

Allows export of currently selected nodes in the Cytoscape window for data export

Methods

signature(descStr = "character", cwObj = "numeric", saveObj = "list") descStr is a string describing the nodes that are currently selected, cwObj is the CytoscapeWindow that the nodes are in, and then saveObj is a previously generated cytOutNodes list, and is optional.

Examples

```
## Not run:
hasCy <- (if (.Platform$OS.type %in% "windows") {
  (length(grep("Cytoscape", system("tasklist", intern=TRUE))) > 0)})

if hasCy {
  ccResults$BP <- breakEdges(ccResults$BP, 0.8)
  cwObj <- ccOutCyt(ccResults$BP, ccOpts)
  # user selects some nodes in Cytoscape
  RCy3::selectNodes(cwObj, c("GO:0007017", "GO:0000226", "GO:0007051", "GO:0007052"))
  savedNodes <- cytOutNodes("random1", cwObj) # save them
  # and selects some other nodes
  RCy3::selectNodes(cwObj,
    c("GO:0071103", "GO:0034728", "GO:0006323", "GO:0030261", "GO:0006334"),
    preserve.current.selection=FALSE)
  savedNodes <- cytOutNodes("random2", cwObj, savedNodes)

}

## End(Not run)
```

fdr *Number of FDR runs to perform*

Description

Queries or sets the number of random runs to perform to generate an estimate of the false discovery rate. Defaults to 50

Usage

fdr(object)

Arguments

object Can be ccGeneList, HyperGParamsCC, HyperGResultCC, ccEnrichResult See Details for more information.

Details

fdr(object) gets the number of fdr runs for ccGeneList, HyperGParamsCC, HyperGResultCC, ccEnrichResult

fdr(object)<- will set the number of fdr runs to be used by ccEnrich and HyperGTestCC when performing calculations on either a ccGeneList or HyperGParamsCC, respectively

Author(s)

Robert M Flight

See Also

[HyperGResultCC](#) [ccEnrichResult](#) [ccGeneList](#) [HyperGParamsCC](#)

GENccEnrichResult-class

Class "GENccEnrichResult"

Description

Holds generic ccEnrich type results

Objects from the Class

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

Slots

.Data: Object of class "list" The actual list containing the ccEnrichResults

categoryName: Object of class "character"

ontology: Object of class "character"

geneAnnMapping: Object of class "namedList"

graphType: Object of class "character"

names: Object of class "character"

Extends

Class "[namedList](#)", directly. Class "[list](#)", by class "namedList", distance 2. Class "[vector](#)", by class "namedList", distance 3. Class "[AssayData](#)", by class "namedList", distance 3.

Methods

```
[ signature(x = "GENccEnrichResult", i = "ANY", j = "ANY"): Subsets the object to just those
  lists that are desired
categoryName signature(object = "GENccEnrichResult"):
ccCompareGeneric signature(gccResult = "GENccEnrichResult", ccOptions = "ccOptions"):
  ...
geneAnnMapping signature(object = "GENccEnrichResult"): ...
graphType signature(object = "GENccEnrichResult"): ...
graphType<- signature(object = "GENccEnrichResult"): ...
ontology signature(object = "GENccEnrichResult"): ...
```

Author(s)

Robert M Flight

See Also

[ccCompareGeneric](#) [ccSigList](#)

Examples

```
data(ccData)
locA <- grep("A", gseaRes$Tissues)
locL <- grep("L", gseaRes$Tissues)
locM <- grep("M", gseaRes$Tissues)

A <- new("ccSigList", sigID=gseaRes$KEGGID[locA], categoryName="KEGG", annotation="org.Mm.eg")
L <- new("ccSigList", sigID=gseaRes$KEGGID[locL], categoryName="KEGG", annotation="org.Mm.eg")
M <- new("ccSigList", sigID=gseaRes$KEGGID[locM], categoryName="KEGG", annotation="org.Mm.eg")
ccEnrichCol <- list(A=A, L=L, M=M)
ccEnrichCol <- new("GENccEnrichResult", ccEnrichCol, categoryName="KEGG")
```

getGeneSymbol	<i>Entrez to name, symbol, GO and path conversion, as well as general ID to ID conversion.</i>
---------------	--

Description

Get different attributes for the Entrez gene Ids

Usage

```
getGeneSymbol(id, annPackage)
getGeneName(id, annPackage)
getG02ALLEGS(id, annPackage)
getPath2EG(id, annPackage)
getAnnotation(id, annPackage, mapID, doUnlist=TRUE)
```

Arguments

id	The IDs one wants to get information for.
annPackage	Which annotation package to use.
mapID	Which mapping to use
doUnlist	should the results be unlisted or not?

Details

The type of ID will change depending on the function. For `getGene...` the ID should be Entrez IDs. For `getG02ALLEGs` Gene Ontology IDs should be used, and for `getPath2EG` KEGG pathways IDs should be used. For `getAnnotation`, any ID can be used.

Value

Returns the requested information.

Note

These functions are generally called internally for mapping between genes and various objects.

Author(s)

Robert M Flight

graphType-methods	<i>graphType</i>
-------------------	------------------

Description

Gets and sets the `graphType` for a couple of different `ccEnrichResults` objects

Methods

```
signature(object = "ccEnrichResult")
signature(object = "GENccEnrichResult")
```

See Also

[ccEnrichResult](#) [GENccEnrichResult](#)

HyperGParamsCC-class *Class "HyperGParamsCC"*

Description

This class extends the HyperGParams class in Category by providing options for multiple testing and the storing of extra data in addition to the gene list of interest (not currently used, but might be in the future).

Objects from the Class

Objects can be created by calls of the form `new("HyperGParamsCC", ...)`. In general the user will not create these directly, but they are created and used by `ccEnrich` to carry out the enrichment calculations.

Slots

fdr: Object of class "numeric" The number of FDR runs to perform
data: Object of class "data.frame" Extra data stored in the object
geneIds: Object of class "ANY" The genes of interest
universeGeneIds: Object of class "ANY" The gene universe or background used (all the genes on the chip)
annotation: Object of class "character" The annotation package used to get information about the geneIds
datPkg: Object of class "DatPkg" Generated automatically from the annotation slot
categorySubsetIds: Object of class "ANY" A specific set of category IDs that one wants to restrict the testing to
categoryName: Object of class "character" What type of category to use, currently either "GO" or "KEGG"
pvalueCutoff: Object of class "numeric" What should be the p-value to decide significance
testDirection: Object of class "character" "over" or "under" represented annotation terms

Extends

Class "[GOHyperGParams](#)", directly.

Methods

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

Author(s)

Robert M Flight

See Also

[HyperGResultCC](#) [ccEnrich](#) Category-package

Examples

```
showClass("HyperGParamsCC")
```

HyperGResultCC-class *Class "HyperGResultCC"*

Description

Contains the results of performing a hypergeometric test on a [HyperGParams](#) object.

Objects from the Class

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

Slots

fdr: Object of class "numeric" The number of FDR runs performed
fdrvalues: Object of class "numeric" The FDR values generated
pvalueType: Object of class "character" Whether to use p-values or FDR values in determining the significant terms returned
data: Object of class "data.frame" Extra data
pvalues: Object of class "numeric" P-values calculated for each term
oddsRatios: Object of class "numeric"
expectedCounts: Object of class "numeric"
catToGeneId: Object of class "list"
organism: Object of class "character"
annotation: Object of class "character"
geneIds: Object of class "ANY"
testName: Object of class "character"
pvalueCutoff: Object of class "numeric"
testDirection: Object of class "character"

Extends

Class "[HyperGResult](#)", directly. Class "[HyperGResultBase](#)", by class "HyperGResult", distance 2.

Methods

fdr signature(object = "HyperGResultCC"): ...
fdr**values** signature(object = "HyperGResultCC"): ...
pCC signature(object = "HyperGResultCC"): ...
pvalueCutoff<- signature(r = "HyperGResultCC"): ...
pvalueType signature(object = "HyperGResultCC"): ...
pvalueType<- signature(object = "HyperGResultCC"): ...
minCount signature(object = "HyperGResultCC"): ...
minCount<- signature(object = "HyperGResultCC"): ...

Author(s)

Robert M Flight

See Also

[hyperGTestCC](#)

Examples

```
showClass("HyperGResultCC")
```

hyperGTestCC

Hypergeometric testing with false discovery rate

Description

Performs the hypergeometric testing for [HyperGParamsCC](#) objects.

Usage

```
hyperGTestCC(p)
```

Arguments

p A [HyperGParamsCC](#) object

Details

This is the heart of categoryCompare, the function that calculates the HyperGeometric statistics for the given categories of annotation for each gene list.

Value

Returns a [HyperGResultCC](#) object

Author(s)

Robert M Flight

See Also

[HyperGParamsCC](#) [HyperGResultCC](#) [GOHyperGParamsCC](#) [KEGGHyperGParamsCC](#) [GOHyperGResultCC](#) [KEGGHyperGResultCC](#)

Examples

```
require(GO.db)
require(org.Hs.eg.db)
data(ccData)
g10 <- unique(table10$Entrez)
testGO <- new("GOHyperGParamsCC", geneIds=g10, universeGeneIds=gUniverse,
  annotation="org.Hs.eg.db", ontology="CC", conditional=FALSE,
  testDirection="over",fdr=0, pvalueCutoff = 0.01)
# ccHypRes <- hyperGTestCC(testGO)
# summary(ccHypRes)
```

<code>listNames</code>	<i>listNames</i>
------------------------	------------------

Description

Extracts the `listNames` from [ccGeneList](#) or [ccOptions](#) objects.

Usage

```
listNames(object)
```

Arguments

`object` This will be either a [ccGeneList](#) or [ccOptions](#) object

Author(s)

Robert M Flight

See Also

[ccGeneList](#) [ccOptions](#)

mergedData-class	Class "mergedData"
------------------	--------------------

Description

Stores merged data tables from the "data" entry in a ccGeneList. This is useful for output later.

Objects from the Class

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

Slots

.Data: Object of class "list"
useIDName: Object of class "character"
names: Object of class "character"
row.names: Object of class "data.frameRowLabels"
.S3Class: Object of class "character"

Extends

Class "[data.frame](#)", directly. Class "[list](#)", by class "data.frame", distance 2. Class "[oldClass](#)", by class "data.frame", distance 2. Class "[data.frameOrNULL](#)", by class "data.frame", distance 2. Class "[vector](#)", by class "data.frame", distance 3.

Methods

`signature(saveObj = "list", compareResult = "ccCompareResult", mergedData = "mergedData")`

Author(s)

Robert M. Flight

See Also

[mergeLists](#) [cytOutData](#)

Examples

```
showClass("mergedData")

data(ccData)
mergeDat <- mergeLists(geneLists, ccOpts)
```

mergeLists-methods *Function mergeLists in Package **categoryCompare***

Description

Merges the gene lists or the data tables from a `ccGeneList` object, providing a single table with all the input data, that can then be queried later, using `cytTableOut`

Usage

```
mergeLists(ccGeneList, ccOptions, isGene=TRUE)
```

Arguments

<code>ccGeneList</code>	a <code>ccGeneList</code> object
<code>ccOptions</code>	a <code>ccOptions</code> object
<code>isGene</code>	are the identifiers genes, or something else (metabolites, etc)

Value

A `mergedData` object which is really just a glorified data frame. If the `ccGeneList` input had a data list, then these are all merged into a single table. Otherwise, it contains just the gene names and which list they were present in.

Methods

```
signature(ccGeneList = "ccGeneList", ccOptions = "ccOptions")
```

See Also

[ccGeneList](#) [ccOptions](#) [mergedData](#)

Examples

```
data(ccData)
g10 <- (unique(table10$Entrez[1:100]))
g48 <- (unique(table48$Entrez[1:100]))

list10 <- list(genes=g10, universe=gUniverse, annotation="org.Hs.eg.db", data=table10[1:100,])
list48 <- list(genes=g48, universe=gUniverse, annotation="org.Hs.eg.db", data=table48[1:100,])

geneLists <- list(T10=list10, T48=list48)
geneLists <- new("ccGeneList", geneLists, ccType=c("BP", "KEGG"))
ccOpts <- new("ccOptions", listNames = names(geneLists))
mergedDat <- mergeLists(geneLists, ccOpts)

list10 <- list(genes=g10, universe=gUniverse, annotation="org.Hs.eg.db")
list48 <- list(genes=g48, universe=gUniverse, annotation="org.Hs.eg.db")
```

```

geneLists <- list(T10=list10, T48=list48)
geneLists <- new("ccGeneList", geneLists, ccType=c("BP", "KEGG"))
ccOpts <- new("ccOptions", listNames = names(geneLists))
mergedDat <- mergeLists(geneLists, ccOpts)

```

minCount

minCount

Description

Extracts and sets the minimum number of genes that an annotation must have to be considered in subsequent steps.

Usage

```
minCount(object)
```

Arguments

object	This will be either a HyperGResultCC, ccEnrichResult, or ccEnrichCollection object. See Details for more information.
--------	---

Details

minCount(object) fetches the set minCount for HyperGResultCC and ccEnrichResult objects

minCount(object)<- will set the minCount for HyperGResultCC objects, and when applied to ccEnrichResult and ccEnrichCollection sets the minCount for all of the contained objects, so be careful if you want to use different minCounts for different results

Author(s)

Robert M Flight

See Also

[HyperGResultCC](#) [ccEnrichResult](#) [ccEnrichCollection](#)

Examples

```

data(ccData)
enrichLists
minCount(enrichLists) <- 5
enrichLists

```

minNodes*Delete nodes with less than a certain number of genes annotated*

Description

Deletes from the graph those annotations with less than a certain number of genes

Usage

```
minNodes(cwObj, cutoff)
```

Arguments

<code>cwObj</code>	a CytoscapeWindowClass object returned from <code>ccOutCyt</code>
<code>cutoff</code>	the minimum number of genes that an annotation must have

Author(s)

Robert M Flight

See Also

CytoscapeWindowClass [ccOutCyt](#)

Examples

```
## Not run:
hasCy <- (if (.Platform$OS.type %in% "windows") {
  (length(grep("Cytoscape", system("tasklist", intern=TRUE))) > 0)})

if hasCy {
  data(ccData)
  ccResults$BP <- breakEdges(ccResults$BP, 0.8)
  cwObj <- ccOutCyt(ccResults$BP, ccOpts)

  minNodes(cwObj, 5)

}
## End(Not run)
```

pvalueType	Type of p-values to return from object
------------	--

Description

Queries or sets the type of p-values to return from objects, either base calculated (pvals) or from fdr calculations (fdr)

Usage

```
pvalueType(object)
```

Arguments

object	Can be HyperGResultCC, ccEnrichResult, ccEnrichCollection. See Details for more information
--------	---

Details

pvalueType(object) gets the type of p-values to be returned from HyperGResultCC and ccEnrichResult objects

pvalueType(object)<- will set the type of p-values to be returned from HyperGResultCC, ccEnrichResult, ccEnrichCollection. Note that for a ccEnrichCollection, the type is changed for all contained ccEnrichResults

Author(s)

Robert M Flight

See Also

[HyperGResultCC](#) [ccEnrichResult](#) [ccEnrichCollection](#)

Examples

```
# pvalueType-Methods
data(ccData)

## Not run: pvalueType(enrichLists) # this returns an error
pvalueType(enrichLists[[1]])
pvalueType(enrichLists[[1]][[1]])

# change the type for one of the results
pvalueType(enrichLists[[1]]) <- 'pval' # Not recommended practice
enrichLists

# change for all of the results
pvalueType(enrichLists) <- 'pval'
enrichLists
```

resetColors-methods *resetColors*

Description

If the color of particular nodes have been modified from the original color scheme in ccOptions, this will reset them

Methods

signature(cwObj = "numeric", ccOpts = "ccOptions") What CytoscapeWindow to apply this to, and what ccOptions to use for the color scheme.

Optional Arguments: Note that optional arguments include node.attribute.name (default is 'fillcolor') and mode (default is 'lookup')

Author(s)

Robert M Flight

See Also

[ccOptions](#) [setNodeColorMapping](#)

show-methods *Methods for Function show in Package 'categoryCompare'*

Description

The show and summary methods for [HyperGResultCC](#) objects generated using [hyperGTestCC](#)

Methods

show, signature(object = "HyperGResultCC")
summary, signature(object = "HyperGResultCC")

Author(s)

Robert M Flight

Examples

```
## Not run:
data(ccData)
show(enrichLists)
summary(enrichLists[[1]][[1]])

## End(Not run)
```

Index

* classes

ccCompareCollection-class, [7](#)
 ccCompareResult-class, [8](#)
 ccEnrichCollection-class, [12](#)
 ccEnrichResult-class, [13](#)
 ccGeneList-class, [14](#)
 ccOptions-class, [16](#)
 ccSigList-class, [18](#)
 GENccEnrichResult-class, [22](#)
 HyperGParamsCC-class, [25](#)
 HyperGResultCC-class, [26](#)
 mergedData-class, [29](#)

* datasets

ccData, [9](#)

* methods

breakEdges-methods, [4](#)
 ccCompareGeneric-methods, [8](#)
 ccOutCyt-methods, [17](#)
 cwReload-methods, [19](#)
 cytOutData-methods, [20](#)
 cytOutNodes-methods, [21](#)
 graphType-methods, [24](#)
 mergeLists-methods, [30](#)
 resetColors-methods, [34](#)
 show-methods, [34](#)

* other possible keyword(s)

breakEdges-methods, [4](#)
 ccCompareGeneric-methods, [8](#)
 cwReload-methods, [19](#)
 cytOutNodes-methods, [21](#)
 graphType-methods, [24](#)
 resetColors-methods, [34](#)

* package

categoryCompare-package, [3](#)

[, GENccEnrichResult, ANY, ANY, ANY-method
 (GENccEnrichResult-class), [22](#)

[, GENccEnrichResult, ANY, ANY-method
 (GENccEnrichResult-class), [22](#)

[, ccEnrichResult, ANY, ANY, ANY-method

(ccEnrichResult-class), [13](#)
 [, ccEnrichResult, ANY, ANY-method
 (ccEnrichResult-class), [13](#)

allAnnotation (ccCompareResult-class), [8](#)

allAnnotation, ccCompareResult-method
 (ccCompareResult-class), [8](#)

annotation, ccSigList-method
 (ccSigList-class), [18](#)

AssayData, [7](#), [12](#), [13](#), [22](#)

breakEdges, [5](#), [7](#), [18](#)

breakEdges (breakEdges-methods), [4](#)

breakEdges, ccCompareResult, numeric-method
 (breakEdges-methods), [4](#)

breakEdges, numeric, numeric-method
 (breakEdges-methods), [4](#)

breakEdges-methods, [4](#)

category, ccSigList-method
 (ccSigList-class), [18](#)

category, GENccEnrichResult-method
 (GENccEnrichResult-class), [22](#)

categoryCompare
 (categoryCompare-package), [3](#)

categoryCompare-package, [3](#)

ccCompare, [7–10](#), [12](#), [18](#)

ccCompare (ccCompare-methods), [5](#)

ccCompare, ccEnrichCollection, ccOptions-method
 (ccCompare-methods), [5](#)

ccCompare, GENccEnrichResult, ccOptions-method
 (ccCompare-methods), [5](#)

ccCompare, G0ccEnrichResult, ccOptions-method
 (ccCompare-methods), [5](#)

ccCompare, KEGGccEnrichResult, ccOptions-method
 (ccCompare-methods), [5](#)

ccCompare-methods, [5](#)

ccCompareCollection, [7](#), [9](#), [10](#)

ccCompareCollection
 (ccCompareCollection-class), [7](#)

- ccCompareCollection-class, 7
- ccCompareGeneric, 19, 23
- ccCompareGeneric
 - (ccCompareGeneric-methods), 8
- ccCompareGeneric, GENccEnrichResult, ccOptions-method
 - (GENccEnrichResult-class), 22
- ccCompareGeneric-methods, 8
- ccCompareResult, 5, 7, 8, 18
- ccCompareResult
 - (ccCompareResult-class), 8
- ccCompareResult-class, 8
- ccData, 9
- ccEnrich, 5, 7, 10, 12, 25
- ccEnrich(ccEnrich-method), 10
- ccEnrich, ccGeneList-method
 - (ccEnrich-method), 10
- ccEnrich-method, 10
- ccEnrichCollection, 10, 31, 33
- ccEnrichCollection
 - (ccEnrichCollection-class), 12
- ccEnrichCollection-class, 12
- ccEnrichResult, 11, 12, 22, 24, 31, 33
- ccEnrichResult(ccEnrichResult-class), 13
- ccEnrichResult-class, 13
- ccGeneList, 10, 11, 22, 28, 30
- ccGeneList(ccGeneList-class), 14
- ccGeneList-class, 14
- ccOptions, 5, 6, 11, 15, 18, 28, 30, 34
- ccOptions(ccOptions-class), 16
- ccOptions, ccCompareGeneric-method
 - (ccCompareGeneric-methods), 8
- ccOptions-class, 16
- ccOpts(ccData), 9
- ccOutCyt, 5, 7, 32
- ccOutCyt(ccOutCyt-methods), 17
- ccOutCyt, ccCompareResult, ccOptions-method
 - (ccOutCyt-methods), 17
- ccOutCyt-methods, 17
- ccResults(ccData), 9
- ccResultsBPHier(ccData), 9
- ccSigList, 23
- ccSigList(ccSigList-class), 18
- ccSigList-class, 18
- ccType(ccGeneList-class), 14
- ccType, ccGeneList-method
 - (ccGeneList-class), 14
- ccType<- (ccGeneList-class), 14
- ccType<-, ccGeneList-method
 - (ccGeneList-class), 14
- compareColors(ccOptions-class), 16
- compareColors, ccOptions-method
 - (ccOptions-class), 16
- compareColors<- (ccOptions-class), 16
- compareColors<-, ccOptions-method
 - (ccOptions-class), 16
- compareIndx(ccOptions-class), 16
- compareIndx, ccOptions-method
 - (ccOptions-class), 16
- compareNames(ccOptions-class), 16
- compareNames, ccOptions-method
 - (ccOptions-class), 16
- compareNames<- (ccOptions-class), 16
- compareNames<-, ccOptions-method
 - (ccOptions-class), 16
- cssClass(ccOptions-class), 16
- cssClass, ccOptions-method
 - (ccOptions-class), 16
- cwReload(cwReload-methods), 19
- cwReload, numeric, ccOptions-method
 - (cwReload-methods), 19
- cwReload, numeric, character, ccOptions-method
 - (cwReload-methods), 19
- cwReload-methods, 19
- cytOutData, 29
- cytOutData(cytOutData-methods), 20
- cytOutData, list, ccCompareResult, mergedData-method
 - (cytOutData-methods), 20
- cytOutData, list, ccCompareResult, missing-method
 - (cytOutData-methods), 20
- cytOutData, list, missing, missing-method
 - (cytOutData-methods), 20
- cytOutData-methods, 20
- cytOutNodes(cytOutNodes-methods), 21
- cytOutNodes, character, numeric, list-method
 - (cytOutNodes-methods), 21
- cytOutNodes, character, numeric, missing-method
 - (cytOutNodes-methods), 21
- cytOutNodes-methods, 21
- data.frame, 29
- data.frameOrNull, 29
- enrichLists(ccData), 9
- fdr, 21

- fdr, ccEnrichResult-method
(ccEnrichResult-class), 13
- fdr, ccGeneList-method
(ccGeneList-class), 14
- fdr, HyperGParamsCC-method
(HyperGParamsCC-class), 25
- fdr, HyperGResultCC-method
(HyperGResultCC-class), 26
- fdr<- (fdr), 21
- fdr<-, ccGeneList-method
(ccGeneList-class), 14
- fdr<-, HyperGParamsCC-method
(HyperGParamsCC-class), 25
- fdrvalues (HyperGResultCC-class), 26
- fdrvalues, HyperGResultCC-method
(HyperGResultCC-class), 26
- fdrvalues-method
(HyperGResultCC-class), 26

- GENccEnrichResult, 19, 24
- GENccEnrichResult
(GENccEnrichResult-class), 22
- GENccEnrichResult, ccCompareGeneric-method
(ccCompareGeneric-methods), 8
- GENccEnrichResult-class, 22
- geneAnnMapping
(GENccEnrichResult-class), 22
- geneAnnMapping, GENccEnrichResult-method
(GENccEnrichResult-class), 22
- geneLists (ccData), 9
- getAnnotation (getGeneSymbol), 23
- getGeneName (getGeneSymbol), 23
- getGeneSymbol, 23
- getGO2ALLEGS (getGeneSymbol), 23
- getPath2EG (getGeneSymbol), 23
- GOccEnrichResult
(ccEnrichResult-class), 13
- GOccEnrichResult-class
(ccEnrichResult-class), 13
- GOHyperGParams, 25
- GOHyperGParams (HyperGParamsCC-class), 25
- GOHyperGParams-class
(HyperGParamsCC-class), 25
- GOHyperGParamsCC, 28
- GOHyperGParamsCC
(HyperGParamsCC-class), 25
- GOHyperGParamsCC-class
(HyperGParamsCC-class), 25

- GOHyperGResultCC, 28
- GOHyperGResultCC
(HyperGResultCC-class), 26
- GOHyperGResultCC-class
(HyperGResultCC-class), 26
- graphType (graphType-methods), 24
- graphType, ccEnrichCollection-method
(ccEnrichCollection-class), 12
- graphType, ccEnrichResult-method
(ccEnrichResult-class), 13
- graphType, GENccEnrichResult-method
(GENccEnrichResult-class), 22
- graphType-methods, 24
- graphType<- (graphType-methods), 24
- graphType<-, ccEnrichCollection-method
(ccEnrichCollection-class), 12
- graphType<-, ccEnrichResult-method
(ccEnrichResult-class), 13
- graphType<-, GENccEnrichResult-method
(GENccEnrichResult-class), 22
- gseaRes (ccData), 9
- gUniverse (ccData), 9

- HyperGParams, 26
- HyperGParamsCC, 22, 27, 28
- HyperGParamsCC (HyperGParamsCC-class), 25
- HyperGParamsCC-class, 25
- HyperGResult, 26
- HyperGResultBase, 26
- HyperGResultCC, 11, 13, 22, 25, 27, 28, 31, 33, 34
- HyperGResultCC (HyperGResultCC-class), 26
- HyperGResultCC-class, 26
- hyperGTestCC, 11, 12, 27, 27, 34
- hyperGTestCC, HyperGParamsCC
(hyperGTestCC), 27
- hyperGTestCC, HyperGParamsCC-method
(hyperGTestCC), 27

- KEGGccEnrichResult
(ccEnrichResult-class), 13
- KEGGccEnrichResult-class
(ccEnrichResult-class), 13
- KEGGHyperGParams
(HyperGParamsCC-class), 25
- KEGGHyperGParams-class
(HyperGParamsCC-class), 25

- KEGGHyperGParamsCC, [28](#)
- KEGGHyperGParamsCC
 - (HyperGParamsCC-class), [25](#)
- KEGGHyperGParamsCC-class
 - (HyperGParamsCC-class), [25](#)
- KEGGHyperGResultCC, [28](#)
- KEGGHyperGResultCC
 - (HyperGResultCC-class), [26](#)
- KEGGHyperGResultCC-class
 - (HyperGResultCC-class), [26](#)
- list, [7](#), [12](#), [13](#), [22](#), [29](#)
- listNames, [28](#)
- listNames, ccGeneList-method
 - (ccGeneList-class), [14](#)
- listNames, ccOptions-method
 - (ccOptions-class), [16](#)
- listNames<-, ccOptions-method
 - (ccOptions-class), [16](#)
- mainGraph (ccCompareResult-class), [8](#)
- mainGraph, ccCompareResult-method
 - (ccCompareResult-class), [8](#)
- mainTable (ccCompareResult-class), [8](#)
- mainTable, ccCompareResult-method
 - (ccCompareResult-class), [8](#)
- mergedData, [30](#)
- mergedData-class, [29](#)
- mergeLists, [29](#)
- mergeLists (mergeLists-methods), [30](#)
- mergeLists, ccGeneList, ccOptions-method
 - (mergeLists-methods), [30](#)
- mergeLists-methods, [30](#)
- minCount, [31](#)
- minCount, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- minCount, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- minCount<- (minCount), [31](#)
- minCount<-, ccEnrichCollection-method
 - (ccEnrichCollection-class), [12](#)
- minCount<-, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- minCount<-, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- minNodes, [32](#)
- minNodes, CytoscapeWindowClass, numeric-method
 - (minNodes), [32](#)
- namedList, [7](#), [12](#), [13](#), [22](#)
- oldClass, [29](#)
- ontology, ccSigList-method
 - (ccSigList-class), [18](#)
- ontology, GENccEnrichResult-method
 - (GENccEnrichResult-class), [22](#)
- outType, [7](#)
- outType (ccOptions-class), [16](#)
- outType, ccOptions-method
 - (ccOptions-class), [16](#)
- outType<- (ccOptions-class), [16](#)
- outType<-, ccOptions-method
 - (ccOptions-class), [16](#)
- pCC, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- pvalueCutoff, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- pvalueCutoff, ccGeneList-method
 - (ccGeneList-class), [14](#)
- pvalueCutoff<-, ccEnrichCollection-method
 - (ccEnrichCollection-class), [12](#)
- pvalueCutoff<-, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- pvalueCutoff<-, ccGeneList-method
 - (ccGeneList-class), [14](#)
- pvalueCutoff<-, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- pvalueType, [33](#)
- pvalueType, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- pvalueType, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- pvalueType<- (pvalueType), [33](#)
- pvalueType<-, ccEnrichCollection-method
 - (ccEnrichCollection-class), [12](#)
- pvalueType<-, ccEnrichResult-method
 - (ccEnrichResult-class), [13](#)
- pvalueType<-, HyperGResultCC-method
 - (HyperGResultCC-class), [26](#)
- resetColors (resetColors-methods), [34](#)
- resetColors, numeric, ccOptions-method
 - (resetColors-methods), [34](#)
- resetColors-methods, [34](#)
- setNodeColorMapping, [34](#)
- show, HyperGResultCC-method
 - (show-methods), [34](#)

show-methods, [34](#)
sigID (GENccEnrichResult-class), [22](#)
sigID, ccSigList-method
 (ccSigList-class), [18](#)
summary, HyperGResultCC-method
 (show-methods), [34](#)
summary-methods (show-methods), [34](#)

table10 (ccData), [9](#)
table48 (ccData), [9](#)
testDirection, ccGeneList-method
 (ccGeneList-class), [14](#)

vector, [7](#), [12](#), [13](#), [22](#), [29](#)