Package 'signet'

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Type Package Title signet: Selection Inference in Gene NETworks Version 1.0.2 Date 2017-11-27 Description An R package to detect selection in biological pathways. Using gene selection scores and biological pathways data, one can search for high-scoring subnetworks of genes within pathways and test their significance. **Depends** R (>= 3.4.0) Imports graph, igraph, RBGL, graphics, utils, stats, methods Suggests graphite, BiocStyle, knitr, rmarkdown biocViews Software, Pathways, DifferentialExpression, GeneExpression, NetworkEnrichment, GraphAndNetwork, KEGG License GPL-2 LazyData true RoxygenNote 6.0.1 VignetteBuilder knitr git_url https://git.bioconductor.org/packages/signet git_branch RELEASE_3_7 git_last_commit f3d0673 git_last_commit_date 2018-06-18 Date/Publication 2018-10-15 Author Alexandre Gouy [aut, cre] Maintainer Alexandre Gouy <alexandre.gouy@iee.unibe.ch>

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kegg_human Pathway examples. See browseVignettes("signet") to see how to prepare the pathways for running simulated annealing.

Description

Pathway examples. See browseVignettes("signet") to see how to prepare the pathways for running simulated annealing.

Usage

kegg_human

Format

An object of class list of length 3.

Null distribution nullDist

Description

Generate the high-scores null distribution to compute empirical p-values for each biological pathway.

Usage

nullDist(pathways, scores, n = 1000, background)

Arguments

pathways	A list of graphNEL objects.
scores	A data frame in which the first column corresponds to the gene ID and the second column contains the gene scores.
n	Number of null high-scores to compute (default = 1000).
background	Optional. Background distribution computed using the backgroundDist func- tion.

Value

A vector of subnetworks scores obtained under the null hypothesis. Must be used as input of the testSubnet function.

scores

Examples

```
# Get KEGG pathways from the package graphite:
# library(graphite)
# kegg <- pathways("hsapiens", "kegg")
# kegg_human <- lapply(kegg, pathwayGraph)
data(daub13) # load the gene scores from Daub et al. (2013)
# generate the null distribution (here, only 5 values, but
# at least 1000 are advised)
null <- nullDist(kegg_human, scores, n = 5)</pre>
```

scores

Example of gene selection scores.

Description

A dataset of gene selection scores.

Usage

scores

Format

A data frame with 17918 rows and 2 variables:

gene Gene identifier (Entrez gene ID)

score Gene selection score ...

Details

These gene scores are the one used in Daub et al. (2013). For a set of SNPs, a z-score has been computed to represent an overall genetic differentiation among several human populations. A gene score corresponds to the maximal value among SNPs located within a given gene.

The table has been generated using the pipeline described here ("Human populations project"): https://github.com/CMPG/polysel

Source

Daub, J. T., Hofer, T., Cutivet, E., Dupanloup, I., Quintana-Murci, L., Robinson-Rechavi, M., & Excoffier, L. (2013). Evidence for polygenic adaptation to pathogens in the human genome. Molecular biology and evolution, 30(7), 1544-1558.

searchSubnet

Description

A simulated annealing algorithm to find the highest scoring subnetwork within a graph.

Usage

```
searchSubnet(pathway, scores, iterations = 1000, background)
```

Arguments

pathway	A gene network, or a list of gene networks, in the graphNEL format.
scores	A data frame with two columns: gene identifiers list (IDs have to be the same as for the pathways, e.g. Entrez) and associated scores.
iterations	Number of iterations.
background	For development purposes.

Value

A signet object or a list of signet objects. Each signet object consists in a table with gene IDs, their state, their score; the subnetwork score and size and the p-value.

Examples

```
# Get KEGG pathways from the package graphite:
# library(graphite)
# kegg <- pathways("hsapiens", "kegg")
# kegg_human <- lapply(kegg, pathwayGraph)
data(daub13) # load the gene scores from Daub et al. (2013)
#run the search in all the pathways with 2500 iterations (default)
example <- searchSubnet(kegg_human, scores)
summary(example)
```

signet

Selection Inference in Gene NETworks

Description

Implements a simulated annealing approach to search for high scoring subnetworks of genes within biological pathways and to test for their significance.

Examples

```
## A complete workflow is described in signet vignette:
browseVignettes("signet")
```

Signet-class

Description

An S4 class to represent a pathway and the results of the associated simulated annealing run.

Usage

```
## S4 method for signature 'Signet'
show(object)
## S4 method for signature 'Signet'
summary(object)
## S4 method for signature 'Signet,missing'
plot(x, y, ...)
## S4 method for signature 'Signet'
initialize(.Object, pathway, scores, iterations)
```

Arguments

object	A signet object.
x	A signet object.
У	Omitted when plotting a Signet object.
	Other graphical parameters.
.Object	Object to initialize.
pathway	Biological pathway (graphNEL object).
scores	Gene scores list.
iterations	Number of simulated annealing iterations.

Value

A plot of the simulated annealing run.

A signet object.

Methods (by generic)

- show: Print the summary a Signet object
- summary: Print the summary of a Signet object
- plot: Plot a Signet object
- initialize: Initialize a Signet object

Slots

connected_comp A graphNEL object (biological pathway)
network A data frame (gene IDs and scores)
SA A data frame (information on the simulated annealing run)
subnet_score A numeric value (subnetwork score)
aggregate_score A numeric value (aggregate subnetwork score)
mean_score A numeric value (average gene score in the pathway)
subnet_size An integer value (subnetwork size)
subnet_genes A factor (subnetwork genes)
p.value A numeric value (empirical p-value)

SignetList-class An S4 class to represent a list of "Signet" objects.

Description

An S4 class to represent a list of "Signet" objects.

Usage

```
## S4 method for signature 'SignetList'
initialize(.Object, list)
## S4 method for signature 'SignetList'
x[[i]]
## S4 method for signature 'SignetList,ANY,missing'
x[i, j, ..., drop = TRUE]
## S4 method for signature 'SignetList'
summary(object, ...)
```

Arguments

.Object	A SignetList object.
list	A list of Signet objects.
x	A SignetList object.
i	Index specifying elements to extract or replace.
j	Unused for SignetList objects.
	Unused for SignetList objects.
drop	Unused for SignetList objects.
object	A SignetList object.

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testSubnet

Value

Results of the simulated annealing run for a lisst of pathways.

A SignetList object.

A data frame containing summary statistics for each element (network and subnetwork sizes, subnetwork score, p-value, significant genes list)

Methods (by generic)

- initialize: Initialize a SignetList
- [[: Access the ith element (signet object) of the SignetList
- [: Access the ith element (signet object) of the SignetList
- summary: Summarize the SignetList.

Slots

results A list of Signet objects.

testSubnet	Test the significance of high-scoring subnetworks found using simu- lated annealing.

Description

Test the significance of high-scoring subnetworks found using simulated annealing.

Usage

```
testSubnet(sigObj, null)
```

Arguments

sigObj	A list of signet objects obtained using the searchSubnet function.
null	Vector of null subnetwork scores generated using the nullDist function.

Value

For each signet object, a p-value is computed given the provided emnpirical null distribution.

Examples

```
# Get KEGG pathways from the package graphite:
# library(graphite)
# kegg <- pathways("hsapiens", "kegg")
# kegg_human <- lapply(kegg, pathwayGraph)
data(daub13) # load the gene scores from Daub et al. (2013)
#run the search in all the pathways with 2500 iterations (default)
example <- searchSubnet(kegg_human, scores)</pre>
```

```
# generate the null distribution (here, only 5 values, but
# at least 1000 are advised)
null <- nullDist(kegg_human, scores, n = 5)
example <- testSubnet(example, null) #now, 'example' includes p-values
summary(example)
```

```
writeXGMML
```

Write Cytoscape input file

Description

This function allows to write an XGMML file to represent the results in Cytoscape.

Usage

```
writeXGMML(sigObj, filename = "signet_output.xgmml", threshold = 0.01)
```

Arguments

sigObj	A signet or signetList object.
filename	The desired file name. Default is "signet_output.xgmml".
threshold	Significance threshold (default: 0.01). If a signetList is provided, all subnetworks with a p-value below this threshold will be merged and represented.

Value

Writes an XGMML file in the working directory. If a single pathway (signet object) is provided, the whole pathway is represented and nodes belonging to the highest-scoring subnetwork (HSS) are highlighted in red. If a list of pathways (signetList) is provided, all subnetworks with a p-value below a given threshold (default: 0.01) are merged and represented. Note that in this case, only the nodes belonging to HSS are kept for representation.

Examples

```
# Get KEGG pathways from the package graphite:
# library(graphite)
# kegg <- pathways("hsapiens", "kegg")
# kegg_human <- lapply(kegg, pathwayGraph)
data(daub13) # load the gene scores from Daub et al. (2013)
#run the search in all the pathways with 2500 iterations (default)
example <- searchSubnet(kegg_human, scores)
#write Cytoscape input file for the first pathway:
writeXGMML(example[[1]], filename = tempfile())
```

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