Package ‘sigora’

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Depends R (>= 2.10)

Imports utils,stats

Suggests slam

Description Pathway Analysis is the process of statistically linking observations on the molecular level to biological processes or pathways on the systems (organism, organ, tissue, cell) level. Traditionally, pathway analysis methods regard pathways as collections of single genes and treat all genes in a pathway as equally informative. This can lead to identification of spurious (misleading) pathways as statistically significant, since components are often shared amongst pathways. SIGORA seeks to avoid this pitfall by focusing on genes or gene-pairs that are (as a combination) specific to a single pathway. In relying on such pathway gene-pair signatures (Pathway-GPS), SIGORA inherently uses the status of other genes in the experimental context to identify the most relevant pathways. The current version allows for pathway analysis of human and mouse data sets and contains pre-computed Pathway-GPS data for pathways in the KEGG and Reactome pathway repositories as well as mechanisms for extracting GPS for user supplied repositories.

License GPL-2

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**R topics documented:**

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**Description**

This package implements the pathway analysis method SIGORA. For an in depth description of the method, please see our manuscript in PeerJ. In short: a GPS (gene pair signature) is a (weighted) pair of genes that as a combination occurs only in a single pathway within a pathway repository. A query list is a vector containing a gene list of interest (e.g. genes that are differentially expressed in a particular condition). A present GPS is a GPS for which both components are in the query list. SIGORA identifies relevant pathways based on the over-representation analysis of their (present) GPS.

**Details**

**Getting started:**
To install from CRAN:
install.packages('sigora')
As an alternative, you can download the tarball and install from the local file:
install.packages("sigora_2.0.1.tar.gz",type="source",repos = NULL)
To load the library:
library("sigora")

**Motivation —A thought experiment:** Imagine you randomly selected 3 KEGG pathways, and then randomly selected a total of 50 genes from all genes that are associated with any of these pathways. Using traditional methods (hypergeometric test using individual genes), how many pathways would you estimate to show up as statistically overrepresented in this "query list" of 50 genes? Let us do this experiment! Everything related to human KEGG Pathways can be found in kegh. A function to randomly select n genes from m randomly selected pathways is genesFromRandomPathways. The traditional Overrepresentation Analysis (which is the basis for
many popular tools) is available through ora. Putting these together:

data(kegH)
a1<-genesFromRandomPathways(seed=12345,kegH,3,50)

## originally selected pathways:
a1["selectedPathways"]
## what are the genes a1["genes"]
## Traditional ora identifies dozens of statistically significant pathways!
ora(a1["genes"],kegH)

## Now let us try sigora with the same input:
sigoraRes <- sigora(GPSrepo =kegH, queryList = a1["genes"],level = 4)
## Again, the three originally selected pathways were:
a1["selectedPathways"]

You might want to rerun the above few lines of code with different values for seed and convince yourself that there indeed is a need for a new way of pathway analysis.

**Available Pathway-GPS repositories in SIGORA:**
The current version of the package comes with precomputed GPS-repositories for KEGG human and mouse (kegH and kegM respectively), as well as for Reactome human and mouse (reaH and reaM respectively). The package provides a function for creating GPS-repositories from user’s own gene-function repository of choice (example Gene Ontology Biological Processes). The following section describes this process of creating one’s own GPS-repositories using the PCI-NCI pathways from National Cancer Institute as an example.

**Creating a GPS repository:** You can create your own GPS repositories using the `makeGPS()` function. There are no particular requirements on the format of your source repository, except: it should be provided either a tab delimited file or a dataframe with **three columns in the following order:**
PathwayID, PathwayName, Gene.

data(nciTable)

## what does the input look like?
head(nciTable)
## create a SigObject. use the saveFile parameter for future reuse.
nciH<-makeGPS(pathwayTable=nciTable, saveFile='nciH.rda')
ils<-grep("^IL",idmap[,"Symbol"],value=TRUE)
ilnci<-sigora(queryList=ils,GPSrepo=nciH,level=3)

**Analysing your data:** To preform Signature Overrepresentation Analysis, use the function `sigora`. For traditional Overrepresentation Analysis, use the function `ora`.

**Exporting the results:** Simply provide a file name to the `saveFile` parameter of `sigora`, i.e. (for the above experiment):
sigRes<- sigora(kegH,queryList= a1$genes,level= 2, saveFile="myResultsKEGG.csv")
You will notice that the file also contains the list of the relevant genes from the query list in each pathway. The genes are listed as human readable gene symbols and sorted by their contribution to the statistical significance of the pathway.
Gene identifier mapping: Mappings between ENSEMBL-IDS, ENTREZ-IDS and Gene-Symbols are performed automatically. You can, for instance, create a GPS-repository using ENSEMBL-IDS and perform Signature Overrepresentation Analysis using this repository on a list of ENTREZ-IDS.

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References
Foroushani AB, Brinkman FS and Lynn DJ (2013). “Pathway-GPS and SIGORA: identifying relevant pathways based on the over-representation of their gene-pair signatures.” PeerJ, 1

See Also
sigora, makeGPS, ora

Examples
barplot(table(kegh$l1$degs),col="red", main="distribution of number of functions per gene in KEGG human pathways.", ylab="frequency",xlab="number of functions per gene")
## creating your own GPS repository
ncih<makeGPS(pathwayTable=nciTable, saveFile='nciH.rda')
ils<-grep("IL",idmap[,"Symbol"],value=TRUE)
## signature overrepresentation analysis:
sigRes.ilnci<-sigora(queryList=ils,GPSrepo=nciH,level=3)

genesFromRandomPathways

Function to randomly select genes associated with randomly pathways.

Description
This function first randomly selects a number (np) of pathways, then randomly selects a number (ng) of genes that are associated with at least one of the selected pathways. The function can be used to compare Sigora’s performance to traditional overrepresentation tests.

Usage
genesisFromRandomPathways(seed = 1234, GPSrepo, np, ng)
getGenes

Arguments

- **seed**
  A random seed.

- **gpsrepo**
  A signature repository (created by ..) or one of the precompiled options.

- **np**
  How many pathways to select.

- **ng**
  Number of genes to be selected.

Value

- **selectedPathways**
  A vector containing the "np" originally selected pathways.

- **genes**
  A vector containing the "ng" selected genes from selectedPathways.

References

Foroushani AB, Brinkman FS and Lynn DJ (2013). “Pathway-GPS and SIGORA: identifying relevant pathways based on the over-representation of their gene-pair signatures.” *PeerJ*, 1

See Also

- [sigora-package](#)

Examples

```r
## select 50 genes from 3 human KEGG pathways
a1<-genesFromRandomPathways(seed=12345,kegH,3,50)
## originally selected pathways:
a1["selectedPathways"]
## what are the genes
a1["genes"]
## sigora's results
sigoraRes <- sigora(GPSrepo = kegH, queryList = a1["genes"],
                    level = 4)
## compare to traditional methods results:
oraRes <- ora(a1["genes"],kegH)
dim(oraRes)
oraRes
```

Description

This function lists the genes involved in the present GPS for a specific pathway in the summary_results.
Usage
getGenes(yy, i)

Arguments
yy A sigora analysis result object (created by sigora).
i The rank position of the pathway of interest in summary_results.

Value
A table (dataframe) with ids of the relevant genes, ranked by their contribution to the statistical significance of the pathway.

See Also
sigora

Examples

```r
a1 <- genesFromRandomPathways(seed=12345, kegH, 3, 50)
## originally selected pathways:
new_a1$selectedPathways
## what are the genes
new_a1$genes
## sigora's results with this input:
sigoraRes <- sigora(GPSrepo = kegH, queryList = new_a1$genes, level = 2)
## Genes related to the second most significant result:
getGenes(sigoraRes, 2)
```

idmap

Identifier mappings for protein coding genes.

Description
A mapping table for ENSEMBL, ENTREZ and gene names (HGNC/MGI symbols) of Human and mouse protein coding gene.

Usage
data("idmap")

Source
www.ensembl.org/biomart/martview

Examples
data(idmap)
head(idmap)
**kegH**

*Pathway GPS data, extracted from KEGG repository (Human)*.

**Description**

KEGG human pathway GPS data, extracted by makeGPS, default settings. This data can be used by sigora to preform signature overrepresenation.

**Usage**

```r
data("kegH")
```

**Source**

<http://www.genome.jp/kegg/pathway.html>

**References**


**See Also**

makeGPS, sigora, reaH

**Examples**

```r
data(kegH)
str(kegH)
```

**kegM**

*Pathway GPS data, extracted from KEGG repository (Mouse)*.

**Description**

KEGG mouse pathway GPS data, extracted by makeGPS, default settings. This data can be used by sigora to preform signature overrepresenation.

**Usage**

```r
data("kegM")
```

**Source**

<http://www.genome.jp/kegg/pathway.html>
References


Examples

```r
data(kegM)
## maybe str(kegM); plot(kegM) ...
```

---

**makeGPS**

*Create your own Signature Object.*

**Description**

Given a repository of gene-pathway associations either in a tab delimited file with three columns (pathwayID, pathway Description, Gene) or a corresponding dataframe, this function identifies all Gene Pair Signatures (pairs of genes that are as a combination unique to a single pathway) and Pathway Unique Genes (genes that are uniquely associated with a single pathway) and stores them in a format that is usable by sigora. Please also see the "details" and "note" sections below.

**Usage**

```r
makeGPS(pathwayTable=NULL, fn=NULL, maxLevels = 5, saveFile=NULL, reponame = "userrepo", maxFunperGene = 100, maxGenesperPathway = 500, minGenesperPathway = 10)
```

**Arguments**

- **pathwayTable** A data frame describing gene-pathway associations in following format: pathwayID, pathwayName, Gene. Either pathwayTable or fn should be provided.
- **fn** Where to find the repository. Either pathwayTable or fn should be provided.
- **maxLevels** For hierarchical repositories, the number of levels to consider.
- **saveFile** Where to save the object as an rda file.
- **reponame** Repository name.
- **maxFunperGene** A cutoff threshold, genes with more than this number of associated pathways are excluded to speed up the GPS identification process.
- **maxGenesperPathway** A cutoff threshold, pathways with more than this number of associated genes are excluded to speed up the GPS identification process.
- **minGenesperPathway** A cutoff threshold, pathways with less than this number of associated genes are excluded to speed up the GPS identification process.
**Details**

The primary purpose of `makeGPS` is to convert a user-supplied gene-pathway association table to a repository of weighted Gene Pair Signatures (GPS) that are unique features of pathways. Such GPS can then be used for signature (gene-pair) based analyses using `sigora`. Additionally, the resulting object also retains the original "single gene"-"pathway" associations for the purpose of followup analyses, such as comparison of `sigora`-results to traditional methods. `ora` is an implementation of the traditional (individual gene) Overrepresentation Analysis.

**Value**

A GPS repository, to be used by `sigora` and `ora`.

**Note**

This function relies on package `slam`, which should be installed from CRAN. It is fairly memory intensive, and it is recommended to be run on a machine with at least 6GB of RAM. Also, make sure to save and reuse the resulting GPS repository in future analyses!

**References**

Foroushani AB, Brinkman FS and Lynn DJ (2013). “Pathway-GPS and SIGORA: identifying relevant pathways based on the over-representation of their gene-pair signatures.” *PeerJ*, 1

**See Also**

`sigora, sigora-package`

**Examples**

```r
data(nciTable)
## what the input looks like:
head(nciTable)
## create a SigObject. use the saveFile parameter for reuse.
nciH<-makeGPS(pathwayTable=nciTable)
ils<-grep("^IL",idmap[,"Symbol"],value=TRUE)
ilnci<-sigora(queryList=ils,GPSrepo=nciH,level=3)
```

---

**nciTable**

*NCI human gene-pathway associations.*

**Description**

PID-NCI human pathway repository, as a data frame with three columns corresponding to: pathwayId, pathwayName, gene. This is an example of the expected format for the input data to `makeGPS`.

**Usage**

```r
data("nciTable")
```
Details

This dataset is provided to illustrate how to create your own GPS repositories. `nciTable` is a dataframe with three columns corresponding to `pathwayId`, `pathwayName` and `gene`. Each row describes the association between an individual gene and a PID-NCI pathway. As you see in the examples section, one can convert this dataframe to a GPS repository using `makeGPS`, and save the results for future reuse. Using the thus created GPS repository one can perform Signature Overrepresentation Analysis on lists of genes of interest.

Source

<https://github.com/NCIP/pathway-interaction-database/tree/master/download>

Examples

```r
data(nciTable)
nCIH <- makeGPS(pathwayTable=nciTable)
data(idmap)
ils <- grep("IL", idmap[, "Symbol"], value=TRUE)
ilnci <- sigora(queryList=ils, GPSrepo=ncIH, level=3)
```

Description

Traditional Overrepresentation Analysis by hypergeometric test: pathways are treated as collections of individual genes and all genes are treated as equally informative. This function is provided for comparison of the results of traditional methods to Sigora.

Usage

```r
ora(genelist, GPSrepo)
```

Arguments

- **genelist**: A vector containing the list of genes of interest (e.g. differentially expressed genes). Following Identifier types are supported: Gene Symbols, ENTREZ-IDs or ENSEMBL-IDs.
- **GPSrepo**: A GPS-repository (either one of the provided precomputed GPS repositories) or one created by `makeGPS`.

Details

The primary purpose of `makeGPS` is to create a GPS repository. It does, however, retain the original "single gene"-"pathway" associations for the purpose of followup analyses, such as comparison of `sigora`-results to traditional methods. `ora` is an implementation of the traditional (individual gene) Overrepresentation Analysis.
**Value**

A dataframe with individual gene ORA results.

**See Also**

`sigora-package`

**Examples**

```r
data(kegmI)
## select 50 genes from 3 mouse pathways
a1<-genesFromRandomPathways(seed=345,kegmI,3,50)
## originally selected pathways:
a1["selectedPathways"]
## compare to traditional methods results:
orares <- ora(a1["genes"],kegmI)
dim(orares)
orares
```

---

**Description**

Reactome human pathway GPS data, extracted by `makeGPS`, default settings. This data can be used by `sigora` to perform signature overrepresentation.

**Usage**

```r
data("reaH")
```

**Source**

[<http://www.reactome.org/>](http://www.reactome.org/)

**References**


**Examples**

```r
data(reaH)
## maybe str(reaH) ; ...
```
Description

Reactome mouse pathway GPS data, extracted by makeGPS, default settings. This data can be used by sigora to preform signature overrepresentation.

Usage

data("reaM")

Source

<http://www.reactome.org/>

References


See Also

makeGPS, sigora, kegm

Examples

data(reaM)
str(reaM)

sigora Sigora’s main function.

Description

This function determines which Signatures (GPS) from a collection of GPS data (GPSrepo argument) for the specified pathway repository are present in the specified list of genes of interest (queryList argument). It then uses the distribution function of hypergeometric probabilities to identify the pathways whose GPS are over-represented among the present GPS and saves the results to the file specified in the saveFile argument.

Usage

sigora(GPSrepo, level, markers = FALSE, queryList = NULL, saveFile=NULL, weighting.method = "invhm")
Arguments

GPSrepo
An object created by makeGPS or one of the precompiled GPS data collections that are provided with this package (currently for KEGG and Reactome). e.g. reaH for human Reactome GPS. See the examples section for creating and using your own GPS.

level
In hierarchical repositories (e.g. Reactome) number of levels to consider. Recommended value for KEGG: 2, for Reactome: 4.

markers
Whether to take single genes that are uniquely associated with only one pathway into account (i.e. should pathway unique genes/PUGs be considered GPS?). Recommended value: TRUE (1).

queryList
A user specified list of genes of interest ("query list"), as a vector of ENSEMBL (recommended)/ENTREZ IDs or gene symbols (HGNC/MGI).

saveFile
If provided, the results are saved here as a tab delimited File (including , for each pathway, a list of genes ordered by their contribution to the statistical significance of the pathway).

weighting.method
The weighting method or GPS. The default weighting scheme for the GPS is the reciproc of the harmonic mean of the degrees of the two component genes of a GPS. A wide range of alternative weighting schemes are pre-implemented (see below). Additional user defined weighting schemes are also supported. Currently, the following alternatives are pre-implemented: 'noweights', 'cosine', 'topov', 'reciprod', 'jac', 'justPUGs' and 'invhm'. Additional user defined weighting schemes are also supported (see section examples).

'noweights': assigns a constant of 1 to all GPS.
'cosine': all GPS are weighted by the cosine of the degrees of their constituent genes.
'topov': all GPS are weighted by topological overlap of their constituent genes.
'reciprod': all GPS are weighted by reciproc of product of the number of pathway annotations of their constituent genes.
'jac': all GPS are weighted by the jaccard similarity of the pathway annotations constituent genes.
'justPUGs': Analysis is performed using PUGs only.
'invhm': all GPS are weighted by the reciproc of the harmonic mean of the degrees of their constituent genes (default).

Value

summary_results
A dataframe listing the analysis results.

detailed_results
A dataframe describing the detailed evidence (present Gene-Pair Signatures) for each pathway.
References

Foroushani AB, Brinkman FS and Lynn DJ (2013). “Pathway-GPS and SIGORA: identifying relevant pathways based on the over-representation of their gene-pair signatures.” PeerJ, 1

See Also

sigora-package, makeGPS

Examples

```r
## query list
ils<-grep("IL", idmap["Symbol"], value=TRUE)
## using precompiled GPS repository:
sigRes.1lreact<-sigora(queryList=ils, GPSrepo=reaH, level=4)
## Not run:
## user created GPS repository:
nciH<-makeGPS(pathwayTable=nciTable)
sigRes.1lnci<-sigora(queryList=ils, GPSrepo=nciH, level=3)
## user defined weighting schemes:
myfunc<-function(a,b){1/log(a+b)}
sigora(queryList=ils, GPSrepo=nciH, level=3, weighting.method = "myfunc")
```

## End(Not run)
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