

Package ‘PubScore’

April 3, 2022

Type Package

Title Automatic calculation of literature relevance of genes

Version 1.6.0

Description

Calculates the importance score for a given gene. The importance score is the total counts of articles in the pubmed database that are a result for that gene AND each term of a list.

Imports ggplot2, igraph, ggrepel, rentrez, progress, graphics, dplyr, utils, methods, intergraph, network, sna

Suggests FCBF, plotly, SummarizedExperiment, SingleCellExperiment, knitr, rmarkdown, testthat (>= 2.1.0), BiocManager, biomaRt

biocViews GeneSetEnrichment, GeneExpression, SystemsBiology, Genetics, Epigenetics, BiomedicalInformatics, Visualization, SingleCell

VignetteBuilder knitr

License MIT + file LICENSE

Encoding UTF-8

LazyData true

RoxygenNote 7.1.1

Depends R (>= 4.0.0)

git_url <https://git.bioconductor.org/packages/PubScore>

git_branch RELEASE_3_14

git_last_commit 9794604

git_last_commit_date 2021-10-26

Date/Publication 2022-04-03

Author Tiago Lubiana [aut, cre],
Helder Nakaya [aut, ths]

Maintainer Tiago Lubiana <tiago.lubiana.alves@usp.br>

R topics documented:

.getSimulation_test	2
.query_pubmed	3
all_counts	3
gene2pubmed_db	4
getScore	4
get_all_counts	5
get_literature_score	6
heatmapViz	7
hgcن_entrez_reference	7
initialize, PubScore-method	8
networkViz	9
plot_literature_graph	9
plot_literature_score	10
pubscore	11
PubScore-class	11
set_all_counts<-	12
test_score	13
Index	15

.getSimulation_test	<i>Auxiliary function for the test method</i>
---------------------	---

Description

Auxiliary function for the test method

Usage

.getSimulation_test(pub, ambiguous = c(), n_simulations)

Arguments

- pub An object of class PubScore
- ambiguous A character vector with possible ambiguous gene names
- n_simulations The number of simulations to run.

Value

A data-frame with a simulation of literature scores for random samplings

<code>.query_pubmed</code>	<code>#' .query_pubmed</code>
----------------------------	-------------------------------

Description

Auxiliary function for getting the list score

Usage

```
.query_pubmed(search_topic, wait_time = 0, ret_max = 1)
```

Arguments

<code>search_topic</code>	Item to search on PubMed via rentrez
<code>wait_time</code>	Time between searches
<code>ret_max</code>	Number of IDs to be returned. Defaults to 1.

Value

The rentrez search result (a list)

<code>all_counts</code>	<code>all_counts</code>
-------------------------	-------------------------

Description

A dataframe with all pubmed counts for the genes in the Dengue dataset in relation to the term "Dengue".

Usage

```
data(all_counts)
```

Format

An object of class `data.frame`

Details

Outcome of the `test_score` method of the `pubscore` class. As this function may take a long time, this dataset speeds up the vignette.

Contains: 3 columns: `#tax_id`: The reference ID for the taxon. All are 9606 (humans). `GeneID`: The Entrez ID code for a given gene. `PubMedID`: A PubMed ID for a paper that mentions the gene in the "Gene ID" column.

1335548 rows: gene-paper associations in the `gene2pubmed` database.

gene2pubmed_db	<i>human genes on gene2pubmed_db</i>
----------------	--------------------------------------

Description

A subset of the gene2pubmed database downloaded via FTP from <ftp://ftp.ncbi.nlm.nih.gov/gene/DATA/gene2pubmed.gz>.
 # The subset contains only the rows corresponding to humans (#tax_id = 906) The table was downloaded in October 2019.

Usage

```
data(gene2pubmed_db)
```

Format

An object of class `data.frame`

Details

Contains: 3 columns: #tax_id: The reference ID for the taxon. All are 9606 (humans). GeneID: The Entrez ID code for a given gene. PubMedID: A PubMed ID for a paper that mentions the gene in the "Gene ID" column.

1335548 rows: gene-paper associations in the gene2pubmed database.

References

Maglott, Donna, et al. 'Entrez Gene: gene-centered information at NCBI.' Nucleic acids research 33.suppl_1 (2005): D54-D58.

getScore	<i>Retrieve the literature_score attribute</i>
----------	--

Description

Retrieve the literature_score attribute

Usage

```
getScore(pub)
```

```
## S4 method for signature 'PubScore'
getScore(pub)
```

Arguments

pub Object of class PubScore

Value

A "numeric" with the literature score for this gene x term combination

Examples

```
# Create a new pubscore object
pub <- pubscore(genes = c('cd4', 'cd8'), terms_of_interest = c('blabla', 'immunity'))
plot(networkViz(pub))
```

get_all_counts	<i>Retrieve the all_counts attribute</i>
----------------	--

Description

Retrieve the all_counts attribute

Usage

```
get_all_counts(pub)

## S4 method for signature 'PubScore'
get_all_counts(pub)
```

Arguments

pub Object of class PubScore

Value

A dataframe containing the counts table for all genes.

Examples

```
# Create a new pubscore object
pub <- pubscore(genes = c('cd4', 'cd8'), terms_of_interest = c('blabla', 'immunity'))
plot(networkViz(pub))
```

```
get_literature_score  get_literature_score
```

Description

Calculates the importance score for a given gene. The importance score is the total counts of articles in the pubmed database that are a result for that gene AND each term of a list

Usage

```
get_literature_score(
  genes,
  terms_of_interest,
  gene2pubmed = FALSE,
  return_all = FALSE,
  wait_time = 0,
  show_progress = TRUE,
  verbose = FALSE
)
```

Arguments

<code>genes</code>	A vector with multiple genes.
<code>terms_of_interest</code>	A list of terms of interest related to the topic you want to find the relevance for
<code>gene2pubmed</code>	logical defining if gene2pubmed db is going to be used. If used, the vector of genes has to be of HUMAN genes in the hgc_n_symbol format.
<code>return_all</code>	Only to be used with gene2pubmed! logical defining if the all_counts table is going to be returned here. Usually it is generated by the test_score function.
<code>wait_time</code>	How long should be the interval between two requests to the ENTREZ database when it fails. Defaults to 0. In seconds.
<code>show_progress</code>	If TRUE, a progress bar is displayed. Defaults to TRUE.
<code>verbose</code>	If TRUE, will display the index of the search occuring. Defaults to FALSE.

Value

A dataframe with the literature scores.

Examples

```
genes <- c('CD8A', 'CD4')
terms_of_interest <-
  c(
    "CD4 T cell",
    "CD14+ Monocyte",
    "B cell",
```

```

      "CD8 T cell",
      "FCGR3A+ Monocyte",
      "NK cell",
      "Dendritic cell",
      "Megakaryocyte",
      'immunity'
    )
  get_literature_score(genes, terms_of_interest)

```

heatmapViz

*Retrieve the heatmap attribute***Description**

Retrieve the heatmap attribute

Usage

```

heatmapViz(pub)

## S4 method for signature 'PubScore'
heatmapViz(pub)

```

Arguments

pub Object of class PubScore

Value

A "gg" object, from ggplot2, containing a heatmap from the counts table.

Examples

```

#Create a new pubscore object
pub <- pubscore(genes = c('cd4', 'cd8'), terms_of_interest = c('blabla', 'immunity'))
plot(heatmapViz(pub))

```

hgcن_entrez_reference *hgcن_entrez_reference*

Description

Contains the result of a query to the biomaRt service done in October, 2019.

Usage

```
data(hgcن_entrez_reference)
```

Format

An object of class `data.frame`

Details

2 columns: `entrezgene_id` (containing the Entrez ids) and `hgnc_symbol` (containing gene symbols from the HUGO gene nomenclature consortium)

20491 rows, for the mapping between the two nomenclatures for human genes.

References

Mapping identifiers for the integration of genomic datasets with the R/Bioconductor package `biomaRt`. Steffen Durinck, Paul T. Spellman, Ewan Birney and Wolfgang Huber, *Nature Protocols* 4, 1184-1191 (2009).

```
initialize, PubScore-method
      initialize
```

Description

`initialize`

Usage

```
## S4 method for signature 'PubScore'
initialize(.Object, genes, terms_of_interest, gene2pubmed = FALSE)
```

Arguments

<code>.Object</code>	The object of signature <code>PubScore</code> that is going to be created
<code>genes</code>	The genes to which you want to calculate and visualize the literature score.
<code>terms_of_interest</code>	A list of terms of interest related to the topic you want to find the relevance.
<code>gene2pubmed</code>	Logical (TRUE / FALSE) defining if <code>gene2pubmed</code> db is going to be used.

Value

A object of the `PubScore` class

networkViz	<i>Retrieve the network attribute</i>
------------	---------------------------------------

Description

Retrieve the network attribute

Usage

```
networkViz(pub)

## S4 method for signature 'PubScore'
networkViz(pub)
```

Arguments

pub Object of class PubScore

Value

A "gg" object, from ggplot2, containing a network from the counts table.

Examples

```
# Create a new pubscore object
pub <- pubscore(genes = c('cd4', 'cd8'), terms_of_interest = c('blabla', 'immunity'))
plot(networkViz(pub))
```

```
plot_literature_graph #'plot_literature_graph
```

Description

Plot a graph inspired in CEMiTool's graphs

Usage

```
plot_literature_graph(
  plot_counts,
  name,
  color = "#B30000FF",
  max_number_of_labels = 10
)
```

Arguments

plot_counts	The dataframe returned from the get_literature_score function
name	The name of the plot.
color	The color of the plot. Defaults to a shade of red ("B30000FF").
max_number_of_labels	The max number of gene labels to show. Defaults to 10.

Value

A plotly/ggplot2 object is either returned or directly plotted

Examples

```
gene <- c('CD4','CD14', "AIF1", "ACVR1", "CDY2A")
terms_of_interest <- c("CD4 T cell", "CD14+ Monocyte")
literature_counts <- get_literature_score(gene, terms_of_interest)
pl <- plot_literature_graph(literature_counts, name = 'test')
pl
```

plot_literature_score *plot_literature_score*

Description

Plots a non-clusterized heatmap of the article counts for the combination of gene list and list of terms NOTE: the object has to be exactly the one returned by get_literature_score.R . Otherwise ggplot2 will not be able to identify the correct columns.

Usage

```
plot_literature_score(plot_counts, return_ggplot = FALSE, is_plotly = FALSE)
```

Arguments

plot_counts	The dataframe returned from the get_literature_score function
return_ggplot	If TRUE, returns a ggplot2 object instead of plotting. Defaults to FALSE.
is_plotly	If TRUE, a interactive plot is plotted in the place o static ggplot. Defaults to FALSE.

Value

A ggplot2 object is either returned or directly plotted

Examples

```
gene <- c('CD4','CD14', "AIF1", "ACVR1", "CDY2A")
terms_of_interest <- c("CD4 T cell", "CD14+ Monocyte", "B cell",
  "CD8 T cell","FCGR3A+ Monocyte", "NK cell", "Dendritic cell",
  "Megakaryocyte", 'immunity')
literature_counts <- get_literature_score(gene, terms_of_interest)
P <-plot_literature_score(literature_counts, return_ggplot = TRUE)
plot(P)
```

pubscore

*PubScore fundamental analysis***Description**

Runs the initialization and the basic functions for querying pubmed and getting the literature scores.

Usage

```
pubscore(terms_of_interest, genes, gene2pubmed = FALSE)
```

Arguments

terms_of_interest	A list of terms of interest related to the topic you want to find the relevance for
genes	A vector with multiple genes.
gene2pubmed	Logical (TRUE / FALSE) defining if gene2pubmed db is going to be used. Defaults to FALSE.

Value

Object of class PubScore

PubScore-class

*An S4 class to represent PubScore results***Description**

The S4 class to PubScore and its basic initialize and show methods.

Slots

terms_of_interest A list of terms of interest related to the topic you want to find the relevance.

genes The genes to which you want to calculate and visualize the literature score.

date The date when the object was initialized. PubScore counts will likely increase with time.

gene2pubmed Logical (TRUE / FALSE) noting if gene to pubmed was used or not.

counts A data frame with the counts retrieved on PubMed

network A visualization of the results found in a network

heatmap A visualization of the results found in a heatmap

set_all_counts<-	<i>Set the all_counts attribute</i>
------------------	-------------------------------------

Description

Set the all_counts attribute

Usage

```
set_all_counts(pub) <- value

## S4 replacement method for signature 'PubScore'
set_all_counts(pub) <- value
```

Arguments

pub	Object of class PubScore
value	The table with all gene x term article counts from the "test_score" method.

Value

A dataframe containing the counts table for all genes.

Examples

```
terms_of_interest <- c('Dengue')
pub <- pubscore(terms_of_interest = terms_of_interest, genes = c("CD4", "CD8", "CD14") )
print(getScore(pub))
data("all_counts")
set_all_counts(pub) <- all_counts
```

test_score

*Test the literature enrichment score***Description**

Test the literature enrichment score

Usage

```
test_score(
  pub,
  total_genes,
  show_progress = TRUE,
  remove_ambiguous = TRUE,
  verbose = FALSE,
  nsim = 1e+05,
  ambiguous_terms = c("PC", "JUN", "IMPACT", "ACHE", "SRI", "SET", "CS", "PROC", "MET",
    "SHE", "CAD", "DDT", "PIGS", "SARS", "REST", "GC", "CP", "STAR", "SI", "GAN", "MARS",
    "SDS", "AGA", "NHS", "CPE", "POR", "MAX", "CAT", "LUM", "ANG", "POLE", "CLOCK",
    "TANK", "ITCH", "SDS", "AES", "CIC", "FST", "CAPS", "COPE", "F2", "AFM", "SPR",
    "PALM", "C2", "BAD", "GPI", "CA2", "SMS", "INVS", "WARS", "HP", "GAL", "SON", "AFM",
    "BORA", "MBP", "MAK", "MALL", "COIL", "CAST ")
)

## S4 method for signature 'PubScore'
test_score(
  pub,
  total_genes,
  show_progress = TRUE,
  remove_ambiguous = TRUE,
  verbose = FALSE,
  nsim = 1e+05,
  ambiguous_terms = c("PC", "JUN", "IMPACT", "ACHE", "SRI", "SET", "CS", "PROC", "MET",
    "SHE", "CAD", "DDT", "PIGS", "SARS", "REST", "GC", "CP", "STAR", "SI", "GAN", "MARS",
    "SDS", "AGA", "NHS", "CPE", "POR", "MAX", "CAT", "LUM", "ANG", "POLE", "CLOCK",
    "TANK", "ITCH", "SDS", "AES", "CIC", "FST", "CAPS", "COPE", "F2", "AFM", "SPR",
    "PALM", "C2", "BAD", "GPI", "CA2", "SMS", "INVS", "WARS", "HP", "GAL", "SON", "AFM",
    "BORA", "MBP", "MAK", "MALL", "COIL", "CAST ")
)
```

Arguments

pub	Object of class PubScore
total_genes	A list of all the possible genes in your study. Usually all the names in the rows of an "exprs" object.
show_progress	If TRUE, a progress bar is displayed. Defaults to True.

<code>remove_ambiguous</code>	If TRUE, ambiguously named genes (such as "MARCH") will be removed. Defaults to TRUE.
<code>verbose</code>	If TRUE, will display the index of the search occurring. Defaults to false.
<code>nsim</code>	The number of simulations to run. Defaults to 100000.
<code>ambiguous_terms</code>	A character vector of the ambiguous terms to use instead of the default. The default includes 60 genes pre-selected as ambiguous (as IMPACT, MARCH and ACHE).

Value

A "gg" object, from ggplot2, containing a network from the counts table.

Examples

```
# Create a new pubscore object
pub <- pubscore(genes = c('cd4', 'cd8'),
  terms_of_interest = c('blabla', 'immunity'))
pub <- test_score(pub,
  total_genes = c('notagene1', 'notagene2', 'cd4', 'cd8'),
  remove_ambiguous = TRUE)
```

Index

- * **datasets**,
 - all_counts, [3](#)
 - gene2pubmed_db, [4](#)
 - hgcn_entrez_reference, [7](#)
- * **literature**
 - all_counts, [3](#)
 - gene2pubmed_db, [4](#)
 - hgcn_entrez_reference, [7](#)
- * **pubmed**,
 - all_counts, [3](#)
 - gene2pubmed_db, [4](#)
 - hgcn_entrez_reference, [7](#)
- * **test**,
 - all_counts, [3](#)
 - gene2pubmed_db, [4](#)
 - hgcn_entrez_reference, [7](#)
- .getSimulation_test, [2](#)
- .query_pubmed, [3](#)
- all_counts, [3](#)
- gene2pubmed_db, [4](#)
- get_all_counts, [5](#)
- get_all_counts, PubScore-method
(get_all_counts), [5](#)
- get_literature_score, [6](#)
- getScore, [4](#)
- getScore, PubScore-method (getScore), [4](#)
- heatmapViz, [7](#)
- heatmapViz, PubScore-method
(heatmapViz), [7](#)
- hgcn_entrez_reference, [7](#)
- initialize, PubScore-method, [8](#)
- networkViz, [9](#)
- networkViz, PubScore-method
(networkViz), [9](#)
- plot_literature_graph, [9](#)
- plot_literature_score, [10](#)
- pubscore, [11](#)
- PubScore-class, [11](#)
- set_all_counts<-, [12](#)
- set_all_counts<-, PubScore-method
(set_all_counts<-), [12](#)
- test_score, [13](#)
- test_score, PubScore-method
(test_score), [13](#)