

Package ‘pageRank’

April 10, 2023

Title Temporal and Multiplex PageRank for Gene Regulatory Network Analysis

Version 1.8.0

Description Implemented temporal PageRank analysis as defined by Rozenshtein and Gionis. Implemented multiplex PageRank as defined by Halu et al. Applied temporal and multiplex PageRank in gene regulatory network analysis.

Depends R (>= 4.0)

License GPL-2

Encoding UTF-8

LazyData true

Imports GenomicRanges, igraph, motifmatchr, stats, utils, grDevices, graphics

Suggests bcellViper, BSgenome.Hsapiens.UCSC.hg19, JASPAR2018, TxDb.Hsapiens.UCSC.hg19.knownGene, org.Hs.eg.db, TFBSTools, GenomicFeatures, annotate

biocViews StatisticalMethod, GeneTarget, Network

URL <https://github.com/hd2326/pageRank>

BugReports <https://github.com/hd2326/pageRank/issues>

RoxygenNote 6.1.99.9001

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

git_branch RELEASE_3_16

git_last_commit b9d6c66

git_last_commit_date 2022-11-01

Date/Publication 2023-04-10

Author Hongxu Ding [aut, cre, ctb, cph]

Maintainer Hongxu Ding <hd2326@columbia.edu>

R topics documented:

accessibility_network	2
adjust_graph	3
aracne_network	4
bubble_plot	5
clean_graph	6
conformation_network	6
diff_graph	8
get_color_gradient	9
multiplex_page_rank	10
P_graph	11
P_null	12
time_expmat	13

Index	14
--------------	-----------

accessibility_network *Build Network from Accessibility Peaks.*

Description

Build network from accessibility, e.g. ATAC-Seq peaks.

Usage

```
accessibility_network(table, promoter, pfm, genome, p.cutoff = 5e-05, w = 7)
```

Arguments

table	(data.frame) Peaks, with "Chr", "Start" and "End" in column name, and peak ID in row names.
promoter	(GRanges) Promoter regions.
pfm	(PFMatrixList) Position Frequency Matrices (PFMs) of regulators.
genome	(BSgenome or character) Genome build in which regulator motifs will be searched.
p.cutoff	(numeric) P-value cutoff for motifs searching within peaks for TF identification.
w	(numeric) Window size for motifs searching within peaks for TF identification.

Value

(data.frame) Network, with "reg" and "target" in column name.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```

table <- data.frame(Chr=c("chr1", "chr1"), Start=c(713689, 856337),
                     End=c(714685, 862152), row.names=c("A", "B"),
                     stringsAsFactors=FALSE)
regulators=c("FOXF2", "MZF1")
#peaks and regulators to be analyzed

library(GenomicRanges)
library(GenomicFeatures)
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(org.Hs.eg.db)
library(annotate)
promoter <- promoters(genes(TxDb.Hsapiens.UCSC.hg19.knownGene))
names(promoter) <- getSYMBOL(names(promoter), data="org.Hs.eg")
promoter <- promoter[!is.na(names(promoter))]
#get promoter regions

library(JASPAR2018)
library(TFBSTools)
library(motifmatchr)
pfm <- getMatrixSet(JASPAR2018, list(species="Homo sapiens"))
pfm <- pfm[unlist(lapply(pfm, function(x) name(x))) %in% regulators]
#get regulator position frequency matrix (PFM) list

library(BSgenome.Hsapiens.UCSC.hg19)
accessibility_network(table, promoter, pfm, "BSgenome.Hsapiens.UCSC.hg19")
#generate network

```

adjust_graph

Re-calculate PageRank

Description

Re-calculate PageRank with updated damping factor, personalized vector and edge weights.

Usage

```
adjust_graph(graph, damping = 0.85, personalized = NULL, weights = NULL)
```

Arguments

- | | |
|--------------|------------------------------------|
| graph | (igraph) The graph to be adjusted. |
| damping | (numeric) Damping factor. |
| personalized | (numeric) Personalized vector. |
| weights | (numeric) Weight vector. |

Value

(igraph) Network with updated "pagerank" as vertex attribute.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
library(igraph)
set.seed(1)
graph <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph)$name <- 1:100
igraph::V(graph)$pagerank <- igraph::page_rank(graph, damping=0.85)$vector
adjust_graph(graph, damping=0.1)
```

aracne_network

Re-format ARACNe Network.

Description

Re-format ARACNe network in regulon object to data.frame with regulator, target and direction columns.

Usage

```
aracne_network(regulon)
```

Arguments

regulon (regulon) ARACNe network.

Value

(data.frame) Network, with "reg", "target" and "direction" in column name. For direction, 1/0 denotes positive/negative regulation.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
library(bcellViper)
data(bcellViper)
aracne_network(regulon[1:10])
```

bubble_plot	<i>Make Bubbleplot</i>
-------------	------------------------

Description

Make bubbleplot.

Usage

```
bubble_plot(  
  s_mat,  
  c_mat,  
  n_mat,  
  col = colorRampPalette(c("Blue", "Grey", "Red"))(100),  
  breaks = seq(-2, 2, length.out = 100),  
  main = NULL  
)
```

Arguments

s_mat	(matrix) Matrix denotes the size of bubbles.
c_mat	(matrix) Matrix denotes the color of bubbles.
n_mat	(matrix) Matrix denotes the name of bubbles.
col	(character) Colors.
breaks	(numeric) Breakpoints of colors.
main	(character) Title.

Value

(NULL) A bubbleplot.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
s_mat <- c_mat <- n_mat <- matrix(1:12, 3, 4, dimnames=list(1:3, 1:4))  
bubble_plot(s_mat, c_mat, n_mat, breaks=seq(1, 12, length.out=100), main="")
```

clean_graph	<i>Clean Graph</i>
-------------	--------------------

Description

Remove graph nodes by residing subgraph sizes, vertex names and PageRank values.

Usage

```
clean_graph(graph, size = NULL, vertices = NULL, pagerank = NULL)
```

Arguments

graph	(igraph) The graph to be cleaned.
size	(numeric) Subgraph size cutoff.
vertices	(character) Vertices to be kept.
pagerank	(numeric) PageRank cutoff.

Value

(igraph) Network updated "pagerank" as vertex attribute.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
library(igraph)
set.seed(1)
graph <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph)$name <- 1:100
igraph::V(graph)$pagerank <- igraph::page_rank(graph)$vector
clean_graph(graph, size=5)
```

conformation_network	<i>Build Network from Conformation Peaks.</i>
----------------------	---

Description

Build network from conformation, e.g. HiChIP records.

Usage

```
conformation_network(
  table,
  promoter,
  pfm,
  genome,
  range = 500,
  p.cutoff = 5e-05,
  w = 7
)
```

Arguments

table	(data.frame) Records, with "Chr1", "Position1", "Strand1", "Chr2", "Position2" and "Strand2" in column name, and record ID in row names.
promoter	(GRanges) Promoter regions.
pfm	(PFMatrixList) Position Frequency Matrices (PFMs) of regulators.
genome	(BSgenome or character) Genome build in which regulator motifs will be searched.
range	(numeric) Search radius from "Position1" and "Position2" for promoters.
p.cutoff	(numeric) P-value cutoff for motifs searching within peaks for TF identification.
w	(numeric) Window size for motifs searching within peaks for TF identification.

Value

(data.frame) Network, with "reg" and "target" in column name.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
table <- data.frame(Chr1=c("chr1", "chr1"),
                     Position1=c(569265, 713603),
                     Strand1=c("+", "+"),
                     Chr2=c("chr4", "chr1"),
                     Position2=c(206628, 715110),
                     Strand2=c("+", "-"),
                     row.names=c("A", "B"), stringsAsFactors=FALSE)
regulators=c("FOXF2", "MZF1")
#peaks and regulators to be analyzed

library(GenomicRanges)
library(GenomicFeatures)
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
library(org.Hs.eg.db)
library(annotation)
promoter <- promoters(genes(TxDb.Hsapiens.UCSC.hg19.knownGene))
```

```

names(promoter) <- getSYMBOL(names(promoter), data="org.Hs.eg")
promoter <- promoter[!is.na(names(promoter))]
#get promoter regions

library(JASPAR2018)
library(TFBSTools)
library(motifmatchr)
pfm <- getMatrixSet(JASPAR2018, list(species="Homo sapiens"))
pfm <- pfm[unlist(lapply(pfm, function(x) name(x))) %in% regulators]
#get regulator position frequency matrix (PFM) list

library(BSgenome.Hsapiens.UCSC.hg19)
conformation_network(table, promoter, pfm, "BSgenome.Hsapiens.UCSC.hg19")

```

diff_graph*Calculate Temporal PageRank from Two Graphs***Description**

Calculate temporal PageRank by changing edges between graph1 and graph2. This is a simplified version of temporal PageRank described by Rozenshtein and Gionis, by only analyzing temporally adjacent graph pairs.

Usage

```
diff_graph(graph1, graph2)
```

Arguments

graph1	(igraph) The 1st graph.
graph2	(igraph) The 2nd graph.

Value

(igraph) Network graph1-graph2 with "moi (mode of interaction)" and "pagerank" as edge and vertex attributes.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

References

Rozenshtein, Polina, and Aristides Gionis. "Temporal pagerank." Joint European Conference on Machine Learning and Knowledge Discovery in Databases. Springer, Cham, 2016.

Examples

```
library(igraph)
set.seed(1)
graph1 <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph1)$name <- 1:100
set.seed(2)
graph2 <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph2)$name <- 1:100
diff_graph(graph1, graph2)
```

get_color_gradient *Generate Color Gradient*

Description

Generate color gradient for, e.g. gene expression.

Usage

```
get_color_gradient(
  x,
  col = colorRampPalette(c("Blue", "Red"))(100),
  breaks = seq(-2, 2, length.out = 100)
)
```

Arguments

x	(numeric) Vector based on which color gradient is generated.
col	(character) Color vector.
breaks	(numeric) A set of breakpoints for the colors. Must be the same length of col.

Value

(character) Colors.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
get_color_gradient(-2:2)
```

multiplex_page_rank *Calculate Multiplex PageRank***Description**

Calculate multiplex PageRank following definition by Halu et al.

Usage

```
multiplex_page_rank(graph, ..., beta = 1, gamma = 1, damping = 0.85)
```

Arguments

graph	(igraph) The base graph with pagerank and name as vertex attributes.
...	(igraph) Supporter graphs with pagerank and name as vertex attributes.
beta	(numeric) Parameters for adjusting supporter graph PageRank values. For the same nodes, PageRank values from different supporter graphs will first be multiplied. The products will then be exponentiate by beta and gamma, as outgoing edge weights and personalizations of the base graph. Four special multiplex PageRank forms are defined by varying (beta, gamma), including additive (0, 1), multiplicative (1, 0), combined (1, 1) and neutral (0, 0).
gamma	(numeric) Parameters for adjusting supporter graph PageRank values. For the same nodes, PageRank values from different supporter graphs will first be multiplied. The products will then be exponentiate by beta and gamma, as outgoing edge weights and personalizations of the base graph. Four special multiplex PageRank forms are defined by varying (beta, gamma), including additive (0, 1), multiplicative (1, 0), combined (1, 1) and neutral (0, 0).
damping	(numeric) Damping factor.

Value

(numeric) Multiplex PageRank values.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

References

Halu, Arda, et al. "Multiplex pagerank." PloS one 8.10 (2013).

Examples

```
library(igraph)
set.seed(1)
graph1 <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph1)$name <- 1:100
igraph::V(graph1)$pagerank <- igraph::page_rank(graph1)$vector
set.seed(2)
graph2 <- igraph::erdos.renyi.game(100, 0.01, directed = TRUE)
igraph::V(graph2)$name <- 1:100
igraph::V(graph2)$pagerank <- igraph::page_rank(graph2)$vector
multiplex_page_rank(graph1, graph2)
```

P_graph

Build Probability-Based Network

Description

Build probability-based regulator-target interaction network.

Usage

```
P_graph(
  expmat,
  net,
  sep = 5,
  method = c("difference", "mi"),
  null = NULL,
  threshold = 0.001
)
```

Arguments

expmat	(matrix) Gene expression matrix.
net	(data.frame) Network, with "reg" and "target" in column name.
sep	(numeric) Number of bins for calculating marginal/joint probability.
method	(character) Method for calculating probability-based distance, either PXY-PXPY ("difference") or mutual information ("mi").
null	(ecdf) Null distribution of probability-based distance. Either from random interactions by P_null function, or all interactions in net.
threshold	(numeric) P-value threshold for filtering interactions in net.

Value

(igraph) Network graph with "pvalue" and "direction", and "pagerank" as edge/vertex attributes.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
library(bcellViper)
data(bcellViper)
dset <- exprs(dset)
net <- do.call(rbind, lapply(1:10, function(i, regulon){
  data.frame(reg=rep(names(regulon)[i], 10),
             target=names(regulon[[i]][[1]])[1:10],
             direction=rep(1, 10),
             stringsAsFactors = FALSE)}, regulon=regulon))
P_graph(dset, net, method="difference", null=NULL, threshold=0.05)
```

P_null

Build Null Distribution of Probability-Based Distance

Description

Build null model for evaluating the significance of interactions by generating random regulator-target pairs.

Usage

```
P_null(expmat, net, n = 10000, sep = 5, method = c("difference", "mi"))
```

Arguments

<code>expmat</code>	(matrix) Gene expression matrix.
<code>net</code>	(data.frame) Network, with "reg" and "target" in column name.
<code>n</code>	(numeric) Number of random pairs.
<code>sep</code>	(numeric) Number of bins for calculating marginal/joint probability.
<code>method</code>	(character) Method for calculating probability-based distance, either PXY-PXPY ("difference") or mutual information ("mi").

Value

(ecdf) ECDF of null distribution.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
library(bcellViper)
data(bcellViper)
dset <- exprs(dset)
net <- do.call(rbind, lapply(1:10, function(i, regulon){
  data.frame(reg=rep(names(regulon)[i], 10),
             target=names(regulon[[i]][[1]])[1:10],
             direction=rep(1, 10),
             stringsAsFactors = FALSE)}, regulon=regulon))
P_null(dset, net, n=100, method="difference")
```

time_expmat

Generate Timewise Average Gene Expression

Description

Generate timewise average gene expression.

Usage

```
time_expmat(time, expmat)
```

Arguments

time	(character) Time-annotation of samples.
expmat	(matrix) Gene expression matrix.

Value

(matrix) Time-wise average gene expression.

Author(s)

DING, HONGXU (hd2326@columbia.edu)

Examples

```
expmat <- matrix(rnorm(90), 10, 9, dimnames=list(LETTERS[1:10], 1:9))
time <- c(rep("T1", 3), rep("T2", 3), rep("T3", 3))
time_expmat(time, expmat)
```

Index

accessibility_network, 2
adjust_graph, 3
aracne_network, 4

bubble_plot, 5

clean_graph, 6
conformation_network, 6

diff_graph, 8

get_color_gradient, 9

multiplex_page_rank, 10

P_graph, 11
P_null, 12

time_expmat, 13