

Package: BNrich (via r-universe)

August 30, 2024

Type Package

Title Pathway Enrichment Analysis Based on Bayesian Network

Version 0.1.1

Maintainer Samaneh Maleknia <maleknias@gmail.com>

Description Maleknia et al. (2020) <[doi:10.1101/2020.01.13.905448](https://doi.org/10.1101/2020.01.13.905448)>. A novel pathway enrichment analysis package based on Bayesian network to investigate the topology features of the pathways. firstly, 187 kyoto encyclopedia of genes and genomes (KEGG) human non-metabolic pathways which their cycles were eliminated by biological approach, enter in analysis as Bayesian network structures. The constructed Bayesian network were optimized by the Least Absolute Shrinkage Selector Operator (lasso) and the parameters were learned based on gene expression data. Finally, the impacted pathways were enriched by Fisher's Exact Test on significant parameters.

biocViews NetworkEnrichment, GeneExpression, Pathways, Bayesian, KEGG

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Encoding UTF-8

LazyData true

RoxygenNote 7.0.2

Imports bnlearn, corpcor, glmnet, graph, stats, utils

Suggests knitr, rmarkdown

VignetteBuilder knitr

URL <https://github.com/Samaneh-Bioinformatics/BNrich>

BugReports <https://github.com/Samaneh-Bioinformatics/BNrich/issues>

Repository <https://samaneh-bioinformatics.r-universe.dev>

RemoteUrl <https://github.com/samaneh-bioinformatics/bnrich>

RemoteRef HEAD

RemoteSha 8b96e2ced412ecfb7545293cd84975cad03a0c3e

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BNrich	<i>Analysis of significant final BNs</i>
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Description

Fisher's exact test applied to PEA on final BNs

Usage

```
BNrich(Ttest_Results, Pathway.id1, PathName_Final, fdr.value = 0.05)
```

Arguments

Ttest_Results	A data frame contains T-test results for all parameters
Pathway.id1	A vector contains modified KEEG pathway IDs
PathName_Final	A data frame contains is IDs and names of KEEG pathways
fdr.value	A numeric threshold to determine significant parameters

Value

A data frame contains fisher test results for any final pathways

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
M1 <- uni_Result$mapkG1
BN <- BN_struct(M1)
data_h1 <- uni_Result$data_h
data_d1 <- uni_Result$data_d
LASSO_Result <- LASSO_BN(BN = BN , data_h = data_h1 , data_d = data_d1)
```

```

BN_h1 <- LASSO_Result$BN_h
BN_d1 <- LASSO_Result$BN_d
esti_result <- esti_par(BN_H = BN_h1, BN_D = BN_d1, data_h = data_h1, data_d = data_d1)
BNs_H <- esti_result$BNs_h
BNs_D <- esti_result$BNs_d
coef_h <- esti_result$coef_h
coef_d <- esti_result$coef_d
var_result <- var_mat(data_h1, coef_h, BNs_H, data_d1, coef_d, BNs_D)
Var_H = var_result$var_mat_Bh
Var_D = var_result$var_mat_Bd
path.id1 <- uni_Result$pathway.id1
Ttest_result <- parm_Ttest(data_h1, coef_h, BNs_H, data_d1, coef_d, BNs_D, Var_H, Var_D, path.id1)
BNrich_result <- BNrich(Ttest_result, path.id1, Path.Name)

```

BN_struct*Construct Bayesian networks structures***Description**

Construct BNs structures using unified signaling pathways

Usage

```
BN_struct(mapkG1)
```

Arguments

mapkG1	A list contains unified signaling pathways
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Value

A list contains Bayesian networks structures

Examples

```

#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
M1 <- uni_Result$mapkG1
BN <- BN_struct(M1)

```

esti_par*Estimate parameters of BNs in control and disease states***Description**

Estimate parameters of BNs in control and disease states

Usage

```
esti_par(BN_H, BN_D, data_h, data_d)
```

Arguments

BN_H	A list contains simplified BNs structures for control objects
BN_D	A list contains simplified BNs structures for disease objects
data_h	A list contains data frames related to control objects for any BN
data_d	A list contains data frames related to disease objects for any BN

Value

A list contains four lists BNs_h, BNs_d, coef_h and coef_d

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
M1 <- uni_Result$mapkG1
BN <- BN_struct(M1)
data_h1 <- uni_Result$data_h
data_d1 <- uni_Result$data_d
LASSO_Result <- LASSO_BN(BN = BN , data_h = data_h1 , data_d = data_d1)
BN_h1 <- LASSO_Result$BN_h
BN_d1 <- LASSO_Result$BN_d
esti_result <- esti_par(BN_H = BN_h1, BN_D = BN_d1, data_h = data_h1, data_d = data_d1)
```

fetch_data_file	<i>Download data file</i>
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Description

Download necessary data file to start BNrich

Usage

```
fetch_data_file(destfile, verbose = FALSE)
```

Arguments

destfile	A directory in user's own computer for save preprocessed data file
verbose	A logical argument to show verbose results

Value

A list contain mapkG, PathName_final and pathway.id. The mapkG is a list contains imported 187 preprocessed signaling pathways, PathName_final is a data.frame includes names and IDs of all 187 pathways and pathway.id is a character vector of pathways IDs

Examples

```
destfile = tempfile("files", fileext = ".rda")
files <- fetch_data_file()
load(destfile)
```

LASSO_BN	<i>LASSO regression</i>
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Description

LASSO regression – second step of simplification of BNs structures

Usage

```
LASSO_BN(BN, data_h, data_d)
```

Arguments

BN	A list of Bayesian networks achieved by BN_struct function
data_h	A list contains data frames related to control objects
data_d	A list contains data frames related to disease objects

Value

A list contains two lists.BN_h and BN_d are simplified BNs

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
M1 <- uni_Result$mapkG1
BN <- BN_struct(M1)
data_h1 <- uni_Result$data_h
data_d1 <- uni_Result$data_d
LASSO_Result <- LASSO_BN(BN = BN , data_h = data_h1 , data_d = data_d1)
```

parm_Ttest

Testing the equality regression coefficients

Description

t-test for equality the correspongong parameters in any BN

Usage

```
parm_Ttest(
  Data_h,
  coef_H,
  BNs_H,
  Data_d,
  coef_D,
  BNs_D,
  Var_mat_Bh,
  Var_mat_Bd,
  Pathway.id1
)
```

Arguments

Data_h	A list contains data frames related to control objects for any BN
coef_H	A list contains parameters of BN_h
BNs_H	A list contains BNs learned by control objects data
Data_d	A list contains data frames related to disease objects for any BN
coef_D	A list contains parameters of BN_d

BNs_D	A list contains BNs learned by disease objects data
Var_mat_Bh	A list contains covariance matrixes for any node of BN_h
Var_mat_Bd	A list contains covariance matrixes for any node of BN_d
Pathway.id1	A vector contains modified KEEG pathway IDs

Value

A data frame contains T-test results for all parameters in final BNs

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
M1 <- uni_Result$mapkG1
BN <- BN_struct(M1)
data_h1 <- uni_Result$data_h
data_d1 <- uni_Result$data_d
LASSO_Result <- LASSO_BN(BN = BN , data_h = data_h1 , data_d = data_d1)
BN_h1 <- LASSO_Result$BN_h
BN_d1 <- LASSO_Result$BN_d
esti_result <- esti_par(BN_H = BN_h1, BN_D = BN_d1, data_h = data_h1, data_d = data_d1)
BNs_H <- esti_result$BNs_h
BNs_D <- esti_result$BNs_d
coef_h <- esti_result$coef_h
coef_d <- esti_result$coef_d
var_result <- var_mat(data_h1, coef_h, BNs_H, data_d1, coef_d, BNs_D)
Var_H = var_result$var_mat_Bh
Var_D = var_result$var_mat_Bd
path.id1 <- uni_Result$pathway.id1
Ttest_result <- parm_Ttest(data_h1, coef_h, BNs_H, data_d1, coef_d, BNs_D, Var_H, Var_D, path.id1)
```

Description

Unifying nodes based imported signaling pathways and GE data

Usage

```
unify_path(dataH, dataD, MapkG, Pathway.id)
```

Arguments

<code>dataH</code>	A data frame contains (healthy) control objects data
<code>dataD</code>	A data frame contains disease objects data
<code>MapkG</code>	A list contains imported 187 signaling pathways
<code>Pathway.id</code>	A vector contains 187 KEEG pathway IDs

Value

A list contain `data_h`, `data_d`, `mapkG1` and `pathway.id1`

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().
#But here you enter a subset of those pathways to see how this package works.
files <- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)
load(files)
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)
load(Data)
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)
```

var_mat

Estimate variance-covariance matrixes for any parameters of BNs

Description

Estimate variance-covariance matrixes for any parameters of

Usage

```
var_mat(Data_h, coef_H, BNs_H, Data_d, coef_D, BNs_D)
```

Arguments

<code>Data_h</code>	A list contains data frames related to control objects for any BN
<code>coef_H</code>	A lists of parameters of BN_h achieved
<code>BNs_H</code>	A list of BNs learned by control objects data
<code>Data_d</code>	A list contains data frames related to disease objects for any BN
<code>coef_D</code>	A lists of parameters of BN_d
<code>BNs_D</code>	A list of BNs learned by disease objects data

Value

A listcontains two lists `var_mat_Bh` and `var_mat_Bd`

Examples

```
#All the 187 preprocessed signaling pathways can be entered in analysis by fetch_data_file().  
#But here you enter a subset of those pathways to see how this package works.  
files<- system.file("extdata", "test_files_to_start.RData", package = "BNrich", mustWork = TRUE)  
load(files)  
Data <- system.file("extdata", "Test_DATA.RData", package = "BNrich", mustWork = TRUE)  
load(Data)  
uni_Result <- unify_path(dataH, dataD, MapkG = sub_mapkG, Pathway.id = path.id)  
M1 <- uni_Result$mapkG1  
BN <- BN_struct(M1)  
data_h1 <- uni_Result$data_h  
data_d1 <- uni_Result$data_d  
LASSO_Result <- LASSO_BN(BN = BN , data_h = data_h1 , data_d = data_d1)  
BN_h1 <- LASSO_Result$BN_h  
BN_d1 <- LASSO_Result$BN_d  
esti_result <- esti_par(BN_H = BN_h1, BN_D = BN_d1, data_h = data_h1, data_d = data_d1)  
BNs_H <- esti_result$BNs_h  
BNs_D <- esti_result$BNs_d  
coef_h <- esti_result$coef_h  
coef_d <- esti_result$coef_d  
var_result <- var_mat(data_h1, coef_h, BNs_H, data_d1, coef_d, BNs_D)
```

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