# Package 'Corbi'

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Version 0.6-2

Title Collection of Rudimentary Bioinformatics Tools

**Description** Provides a bundle of basic and fundamental bioinformatics tools, such as network querying and alignment, subnetwork extraction and search, network biomarker identification.

ByteCompile TRUE

**Depends** R (>= 3.0.2)

Imports Matrix, MASS, stats, CRF, igraph

**Suggests** knitr, rmarkdown, BiocParallel, matrixcalc, mpmi, fitdistrplus

VignetteBuilder knitr

**License** GPL (>= 2)

BugReports https://github.com/wulingyun/Corbi/issues

URL https://github.com/wulingyun/Corbi

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### Description

This pakcage provides a bundle of basic and fundamental bioinformatics tools.

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#### **Details**

These bioinformatics tools are developed by WuLab at Academy of Mathematics and Systems Science, Chinese Academy of Sciences.

Network querying and alignment:

- net\_query Network querying method based on conditional random fields
- net\_query\_batch Batch processing version of net\_query
- net\_align Network alignment method based on conditional random fields

#### Subnetwork extraction and search:

- get\_subnets Enumerate all subnetworks of limited size
- extend\_subnets Extend subnetworks from smaller subnetworks
- best\_subnets Search best subnetworks that maximize given objective function

#### Biomarker identification:

 markrank Biomarker identification and prioritization by integrating gene expression with biomolecular network

#### Differential expression analysis:

• netDEG Sample specific differential expression analysis

#### Data normalization:

• URG\_getFactor Gene expression data normalization by the uniform ratio graph method

#### References

Qiang Huang, Ling-Yun Wu, and Xiang-Sun Zhang. An Efficient Network Querying Method Based on Conditional Random Fields. Bioinformatics, 27(22):3173-3178, 2011.

Qiang Huang, Ling-Yun Wu, and Xiang-Sun Zhang. Corbi: A new R package for biological network alignment and querying. BMC Systems Biology, 7(Suppl 2):S6, 2013.

Duanchen Sun, Xianwen Ren, Eszter Ari, Tamas Korcsmaros, Peter Csermely, and Ling-Yun Wu. Discovering cooperative biomarkers for heterogeneous complex disease diagnoses. Briefings in Bioinformatics, 20(1), 89–101, 2019.

Xinhan Ye, Ling-Yun Wu. URG: a new normalization method for gene expression data based on graph model. Manuscript.

best\_subnets

### **Description**

Search best subnetworks that maximize given objective functions.

### Usage

```
best_subnets(
  func,
  net.matrix,
  max.size = 10,
  exhaust.size = 5,
  max.top = 10000
)
```

### **Arguments**

func The objective function to maximize

net.matrix The adjacent matrix of network

max.size The maximal size of subnetworks

exhaust.size The maximal size of subnetworks that use exhaustive searching strategy

max.top The maiximal number of top candidates kept for evaluation of next size, used in heuristic searching strategy

#### **Details**

Enumerate and search the best subnetworks that maximize given objective function. If the size of subnetworks <= exhaust.size, exact exhaustive searching is applied, otherwise, heuristic searching algorithm is used.

#### Value

A list with the following two components:

subnets The list of top subnetworks in different sizes
obj.values The list of objective values of corresponding subnetworks

#### See Also

```
get_subnets, extend_subnets
```

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#### **Examples**

```
library(Corbi)
net <- matrix(FALSE, nrow=10, ncol=10)
net[sample.int(100, 20)] <- TRUE
net <- net | t(net)
func <- function(subnet) max(subnet) - min(subnet)
result <- best_subnets(func, net, 5)</pre>
```

column

Extract a column from a matrix

### Description

Extract a specified column from a sparse matrix rapidly

### Usage

```
column(m, i)
```

### **Arguments**

m The matrix

i The column index

### **Details**

This function implements faster column extraction algorithm for the CsparseMatrix class in the package **Matrix**.

#### Value

This function will return the specified column as a vector of corresponding type.

extend\_subnets

Extend subnetworks from smaller subnetworks

### **Description**

Extend subnetworks by pairwise overlapping two sets of smaller subnetworks.

```
extend_subnets(subnet1, subnet2, size = 0)
```

#### **Arguments**

subnet1	The matrix representing the first set of subnetworks
subnet2	The matrix representing the second set of subnetworks
size	The desired size of extended subnetworks

#### **Details**

Enumerate all possible subnetworks of desired size by pairwise overlapping two sets of subnetworks of size s1 and s2. The desired size should be between max(s1,s2)+1 and s1+s2-1. Invalid desired size will be replaced by the minimum allowed value max(s1,s2)+1.

### Value

A matrix represents the extended subnetworks, in which each row represents a subnetwork.

### **Examples**

```
library(Corbi)
net <- matrix(FALSE, nrow=10, ncol=10)
net[sample.int(100, 20)] <- TRUE
net <- net | t(net)
subnets <- get_subnets(net, 3)
subnets[[4]] <- extend_subnets(subnets[[3]], subnets[[2]], 4)</pre>
```

get\_adjusted\_deg\_diff Calculate adjusted degree differences for given network

### **Description**

Calculate the adjusted degree differences for all genes in the given network.

#### Usage

```
get_adjusted_deg_diff(net, log.expr.val, scale.degree = FALSE, p = 0.5)
```

### Arguments

net	The binary adjacent matrix of differential expression ratio network.
log.expr.val	Numeric vector containing the logarithmic scale gene expression values.
scale.degree	Logical variable indicating whether the degree values are scaled according to the dropout rate.
р	The parameter for calculating the adjusted degree differences.

get\_diff\_ratio\_net 7

### Value

This function will return a list with the following components:

diff A numeric vector containing the adjusted degree differences of all genes.

degree A list containing the raw degree differences and sums of all genes.

### **Description**

Construct the differential expression ratio network for a single sample.

### Usage

```
get_diff_ratio_net(
  ref.ratio.dist,
  expr.val,
  log.expr = FALSE,
  scale.degree = FALSE
)
```

### Arguments

ref.ratio.dist	The expression ratio distribution profile returned by $get\_ratio\_distribution$ or $get\_ratio\_distribution2$ .
expr.val	Numeric vector of gene expression values in the sample.
log.expr	Logical variable indicating whether the input expression vector is in logarithmic scale.
scale.degree	Logical variable indicating whether the degree values are scaled according to the dropout rate.

### Value

This function will return a list with the following components:

net The binary adjacent matrix of differential expression ratio network.

diff A numeric vector containing the adjusted degree differences of all genes.

degree A list containing the raw degree differences and sums of all genes.

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```
get_ratio_distribution
```

Calculate expression ratio distribution

#### **Description**

Calculate the lower and upper quantiles of expression ratios for each pair of genes, and estimate the parameters of negative binomial distribution from reference expression data.

### Usage

```
get_ratio_distribution(
  ref.expr.matrix,
  p.edge = 0.1,
  log.expr = FALSE,
  scale.degree = FALSE,
  use.parallel = FALSE
)
```

### Arguments

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The reference expression matrix. Each row represents a gene and each column

represents a sample.

p.edge The expected probability of edges in the expression ratio network for a normal

sample.

log.expr Logical variable indicating whether the input expression matrix is in logarithmic

scale.

scale.degree Logical variable indicating whether the degree values are scaled according to

the dropout rate.

use.parallel Logical variable indicating to use the BiocParallel package to accelerate com-

putation.

### Value

This function will return a list with the following components:

LB	A numeric matrix	with element	ſi.i	l represents the	lower o	quantile of expre	essioin

ratios for gene pairs (i, j).

NB A numeric vector with two elements: size and mu, which are the estimated

parameters of negative binomial distribution.

p.edge The used input parameter p.edge.

get\_ratio\_distribution2 9

```
get_ratio_distribution2
```

Calculate expression ratio distribution

### Description

Calculate the lower and upper quantiles of expression ratios after trimming the extreme values, and estimate the parameters of negative binomial distribution from reference expression data.

### Usage

```
get_ratio_distribution2(
  ref.expr.matrix,
  p.edge = 0.1,
  p.trim = 0.3,
  log.expr = FALSE,
  scale.degree = FALSE,
  use.parallel = FALSE
)
```

### **Arguments**

ref.expr.matri	x
	The reference expression matrix. Each row represents a gene and each column represents a sample.
p.edge	The expected probability of edges in the expression ratio network for a normal sample.
p.trim	The percentage of lower or upper extreme values to be trimmed from the expression ratios for each pair of genes.
log.expr	Logical variable indicating whether the input expression matrix is in logarithmic scale.
scale.degree	Logical variable indicating whether the degree values are scaled according to the dropout rate.
use.parallel	Logical variable indicating to use the BiocParallel package to accelerate computation.

### Value

This function will return a list with the following components:

LB	A numeric matrix with element [i,j] represents the lower quantile of trimmed expressioin ratios for gene pairs (i, j).
NB	A numeric vector with two elements: size and mu, which are the estimated parameters of negative binomial distribution.
p.edge	The used input parameter p.edge.
p.trim	The used input parameter p.trim.

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Calculate expression ratio variances

### Description

Calculate the variances of expression ratios for each pair of genes.

#### Usage

```
get_ratio_variance(expr.matrix, log.expr = FALSE)
```

### Arguments

expr.matrix The expression matrix. Each row represents a gene and each column represents

a sample.

log.expr Logical variable indicating whether the input expression matrix is in logarithmic

scale.

#### Value

This function will return a numeric matrix with element [i,j] represents the variance of expression ratios for gene pairs (i, j).

```
get_shortest_distances
```

Calculate shortest distances of unweighted network

#### **Description**

Calculate all pairs of shortest distances of unweighted network

### Usage

```
get_shortest_distances(
  net.matrix,
  source.nodes = rep_len(TRUE, dim(net.matrix)[1])
)
```

#### **Arguments**

net.matrix Logical adjacency matrix of given unweighted network

distances

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#### **Details**

This function calculates all pairs of shortest distances of unweighted network by using breadth-first-search (BFS) algorithm.

#### Value

This function will return the shortest distance matrix, where the element [i, j] is the shortest distance between node i and j. Value -1 means unreachable. If source.nodes[i] equals FALSE, the shortest distance from i to other nodes will not be calculated and the row i will be all -1.

get\_subnets

All subnetworks of limited size

### **Description**

Enumerate all subnetworks of size <= max.size from given network.

### Usage

```
get_subnets(net.matrix, max.size = 2)
```

### Arguments

net.matrix The adjacent matrix of network
max.size The maximal size of subnetworks

### Value

A list of generated subnetworks, with element \$i\$ corresponds the subnetworks of size \$i\$. Each element is a matrix, in which each row represents a subnetwork.

### **Examples**

```
library(Corbi)
net <- matrix(FALSE, nrow=10, ncol=10)
net[sample.int(100, 20)] <- TRUE
net <- net | t(net)
subnets <- get_subnets(net, 3)</pre>
```

make\_DEG\_data

kappa\_score

Cohen's kappa score

### Description

Calculate Cohen's kappa score for two vectors.

### Usage

```
kappa_score(x1, x2)
```

### Arguments

x1 The first logical vectorx2 The second logical vector

### **Details**

This function calculate Cohen's kappa score for two logical vectors.

### Value

The Cohen's kappa score

make\_DEG\_data

Simulate differentially expressed gene data (Gaussian)

### **Description**

Generate differentially expressed gene (DEG) data from Gaussian distribution.

```
make_DEG_data(
    n.genes,
    n.samples.A,
    n.samples.B,
    exp.mean = 8,
    exp.sd = 2,
    alpha = 0.2,
    size.factor.sd = 0.1,
    ...
)
```

make\_DEG\_data2

#### **Arguments**

n.genes	The total number of genes in the simulated data.
n.samples.A	The number of samples in the group A.
n.samples.B	The number of samples in the group B.
exp.mean	The mean of log-normal distribution that determines gene-specific expression mean.
exp.sd	The standard deviation of log-normal distribution that determines gene-specific expression means.
alpha	The dispersion ratio of gene-specific expression standard deviation to mean.
size.factor.sd	The standard deviation of size factors for samples.
	The parameters passed to function make_DEG_pattern.

### **Details**

The expression values of each gene are assumed following a Gaussian distribution with genespecific mean, which follows a log-normal distribution. The size factor for each sample follows a Gaussian distribution with zero mean and specific standard deviation. The heterogeneity of gene expression data is simulated by using the function make\_DEG\_pattern.

### Value

This function will return a list with the following components:

DEG	The matrix of simulated DEG pattern, which is generated by make_DEG_pattern.
countsA	The expression matrix of group A. Each row represents a gene and each column represents a sample.
countsB	The expression matrix of group B. Each row represents a gene and each column represents a sample.

make_DEG_data2	Simulate differentially expressed gene data (Negative binomial)

### Description

Generate differentially expressed gene (DEG) data from negative binomial distribution.

```
make_DEG_data2(
    n.genes,
    n.samples.A,
    n.samples.B,
    exp.mean = 8,
    exp.sd = 2,
```

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```
dispersion = NULL,
size.factor.sd = 0.1,
...
)
```

#### **Arguments**

The total number of genes in the simulated data. n.genes The number of samples in the group A. n.samples.A n.samples.B The number of samples in the group B. exp.mean The mean of log-normal distribution that determines gene-specific expression exp.sd The standard deviation of log-normal distribution that determines gene-specific expression mean. dispersion The dispersion parameter for negative binomial distribution. The default values are determined by the expression mean. size.factor.sd The standard deviation of size factors for samples. The parameters passed to function make\_DEG\_pattern.

#### **Details**

The expression values of each gene are assumed following a negative binomial distribution with gene-specific mean, which follows a log-normal distribution. The size factor for each sample follows a Gaussian distribution with zero mean and specific standard deviation. The heterogeneity of gene expression data is simulated by using the function make\_DEG\_pattern.

### Value

This function will return a list with the following components:

DEG	The matrix of simulated DEG pattern, which is generated by make_DEG_pattern.
countsA	The expression matrix of group A. Each row represents a gene and each column represents a sample.
countsB	The expression matrix of group B. Each row represents a gene and each column represents a sample.

make_DEG_pattern	Simulate differentially expressed gene pattern	
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### Description

Generate complicated differentially expressed gene (DEG) pattern to simulate varied degree of heterogeneity.

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### Usage

```
make_DEG_pattern(
    n.genes,
    n.samples,
    fold.change = 2,
    gene.rate = 0.3,
    sample.rate = 1,
    active.rate = 1,
    up.rate = 0.5
)
```

### Arguments

n.genes	The total number of genes in the simulated data.	
n.samples	The total number of samples in the simulated data.	
fold.change	The fold change level of DEGs.	
gene.rate	The proportion of DEGs to all genes.	
sample.rate	The proportion of abnormal samples to all samples.	
active.rate	The probability that a DEG is truely differentially expressed in an abnormal sample.	
up.rate	The proportion of up-regulated DEGs to all DEGs.	

### Details

The heterogeneity of gene expression pattern is mainly controlled by two parameters: sample.rate and active.rate. If both parameters are equal to 1, the gene expression pattern will be homogeneous, otherwise heterogeneous.

### Value

This function will return a list with the following components:

FC	The matrix of simulated fold changes. Each row represents a gene and each column represents a sample.
gene	The vector of gene status: 1 for up-regulated, -1 for down-regulated, and 0 for normal genes.
sample	The vector of sample status: 1 for abnormal, and 0 for normal samples.

16 markrank

### Description

MarkRank is a novel proposed network-based model, which can identify the cooperative biomarkers for heterogeneous complex disease diagnoses.

### Usage

```
markrank(
  dataset,
  label,
  adj_matrix,
  alpha = 0.8,
  lambda = 0.2,
  eps = 1e-10,
  E_value = NULL,
  trace = TRUE,
  d = Inf,
  Given_NET2 = NULL
)
```

### **Arguments**

dataset	The microarray expression matrix of related disease. Each row represents a sample and each column represents a gene.
label	The 0-1 binary phenotype vector of dataset samples. The size of label must accord with the sample number in dataset.
adj_matrix	The 0-1 binary adjacent matrix of a connected biological network. Here the node set should be the same order as the gene set in expression matrix.
alpha	The convex combination coefficient of network effect and prior information vector E_value. The range of alpha is in [0,1]. A larger alpha will lay more emphasis on the network information. The default value is 0.8.
lambda	In the random walk-based iteration, matrix A1 reflects the stucture information of the biological network, whereas matrix A2 reflects the cooperative effect of gene combinations. Parameter lambda is the convex combination coefficient of two network effects. The range of lambda is in [0,1]. A larger lambda will lay more emphasis on the A1. The default value is 0.2.
eps	The stop criteria for the iterative solution method. The default value is 1e-10.
E_value	A vector containing the prior information about the importance of nodes. Default is the absolute Pearson correlation coefficient (PCC).
trace	Locaical variable indicated whether tracing information on the progress of the gene cooperation network construction is produced.

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d Threshold for simplifying the G\_2 computation. Only the gene pairs whose

shortest distances in PPI network are less than d participate in the G\_2 compu-

tation. The default value is Inf.

Given\_NET2 Whether a computed cooperation network is given for tuning parameter. See

Details for a more specific description.

#### **Details**

MarkRank is a network-based biomarker identification method to prioritize disease genes by integrating multi-source information including the biological network, e.g protein-protein interaction (PPI) network, the prior information about related diseases, and the discriminative power of cooperative gene combinations. MarkRank shows that explicit modeling of gene cooperative effects can greatly improve biomarker identification for complex disease, especially for diseases with high heterogeneity.

MarkRank algorithm contains mainly two steps: 1) The construction of gene cooperation network G\_2 and 2) a random walk based iteration procedure. The following descriptions will help the users to using markrank more convenient:

- 1) As for the construction of the gene cooperation network, we suggest the user to set trace=TRUE to output the G\_2 computation process. The G\_2 construction step finished if the output number is identical to the gene number of the input expression matrix. The parameter d introduced the structure information of used biological network to facilitate the construction of G\_2, only the gene pairs whose shortest distances in network are less than d participate the G\_2 computation. We suggest d=Inf, the default value, to fully use the information of expression matrix. If the user given a preset d, the distance matrix of input network dis will be returned.
- 2) MarkRank uses a random-walk based iteration procedure to score each gene. The detailed formula is:

```
score = alpha*[lambda*A1 + (1-lambda)*A2]*score + (1-alpha)*E_value.
```

The users could set an appropriate parameter settings in their practical application. Our suggested value is alpha=0.8 and lambda=0.2. The model input parameter combinations and iteration steps will be returned in output components initial\_pars and steps, respectively. Because the iteration step is separate with the cooperation network construction, the user can use the parameter Given\_NET2 to tune the model parameters. In detail, the user could set

```
Given_NET2 = result$NET2
```

in markrank input to avoid the repeated computation of G\_2, where the object result is the returned variable of markrank function.

3) The final MarkRank score for each gene is in output score. The users could sort this result and use the top ranked genes for further analysis.

#### Value

This function will return a list with the following components:

score The vector of final MarkRank scores for each gene.

Steps The final iteration steps in random walk based scoring procedure.

NET2 The weighted adjacent matrix of gene cooperation network.

initial\_pars The initial/input parameter values used in MarkRank.

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dis

The pairwise distance matrix of input network. This variable will be Null if input d=Inf.

#### References

Duanchen Sun, Xianwen Ren, Eszter Ari, Tamas Korcsmaros, Peter Csermely, Ling-Yun Wu. Discovering cooperative biomarkers for heterogeneous complex disease diagnoses. Briefings in Bioinformatics, 20(1), 89–101, 2019.

netDEG

netDEG: Differentially expressed gene identification method

#### **Description**

Perform netDEG for two group samples.

### Usage

```
netDEG(
  ref.expr.matrix,
  expr.matrix,
  p.edge = 0.1,
  summarize = c("gene", "sample"),
  summarize.method = c("sumlog", "sumlog"),
  summarize.shrink = c(Inf, Inf),
  log.expr = FALSE,
  zero.as.dropout = TRUE,
  scale.degree = TRUE,
  use.parallel = FALSE
)
```

#### **Arguments**

ref.expr.matrix

The reference expression matrix. Each row represents a gene and each column represents a sample.

expr.matrix The test expression matrix. Each row represents a gene and each column repre-

sents a sample.

p.edge The expected probability of edges in the expression ratio network for a normal

sample.

summarize Character vector indicating how to summarize the results. Available methods

are c("gene", "sample").

summarize.method

Character vector indicating the methods used to summarize the results. See  $p\_combine$ .

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summarize.shrink

Numeric vector indicating the shrink parameter to summarize the results. See

p\_combine.

log.expr Logical variable indicating whether the input expression matrix is in logarithmic

scale.

zero.as.dropout

Logical variable indicating whether the zero expressions are regarded as dropouts.

scale.degree Logical variable indicating whether the degree values are scaled according to

the dropout rate.

use.parallel Logical variable indicating to use the BiocParallel package to accelerate com-

putation.

#### Value

This function will return a list with the following components:

up A numeric matrix with same dimension as expr.matrix, containing the p-

values of up-regulation test.

down A numeric matrix with same dimension as expr.matrix, containing the p-

values of down-regulation test.

twoside A numeric matrix with same dimension as expr.matrix, containing the p-

values of twoside test.

rev A list containing the reverse comparison results, containing three components:

up, down, and twoside. Available if the gene method is specified in summarize

argument.

gene A list containing the gene-wise summaried results, containing three compo-

nents: up, down, and twoside. Available if the gene method is specified in

summarize argument.

sample A list containing the sample-wise summaried results, containing three compo-

nents: up, down, and twoside. Available if the sample method is specified in

summarize argument.

netDEG\_pvalue Calculate netDEG p-values

### Description

Perform the single or two side tests and calculate the p-values.

```
netDEG_pvalue(ref.ratio.dist, expr.val, log.expr = FALSE, scale.degree = FALSE)
```

20 net\_align

### Arguments

ref.ratio.dist	The expression ratio distribution profile returned by get_ratio_distribution or get_ratio_distribution2.
expr.val	Numeric vector of gene expression values in the sample.
log.expr	Logical variable indicating whether the input expression vector is in logarithmic scale.
scale.degree	Logical variable indicating whether the degree values are scaled according to the dropout rate.

#### Value

This function will return a list with the following components:

up A numeric vector containing the p-values of up-regulation test.

down A numeric vector containing the p-values of down-regulation test.

twoside A numeric vector containing the p-values of twoside test.

 ${\tt net\_align} \qquad \qquad {\tt Network~alignment~method~based~on~conditional~random~fields}$ 

### **Description**

Find the maximal matching subnetworks from a target network for a query network based on the conditional random fields (CRF) model.

### Usage

```
net_align(
  query.net,
  target.net,
  node.sim,
  query.type = 4,
  delta.d = 1e-10,
  delta.c = 0.5,
  delta.e = 1,
  delta.s = 1,
  output = "result.txt"
)
```

### Arguments

query.net The input file name of the query network.

target.net The input file name of the target network.

node.sim The input file name of the node similarity scores between the query network and

the target network.

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query.type	The querying network type: 1 - general, 2 - chain, 3 - tree, 4 - heuristic.
delta.d	The parameter delta.d is a parameter for deletions.
delta.c	The parameter delta.c is a parameter for consecutive deletions.
delta.e	The parameter delta.e is a parameter for single deletion.
delta.s	The parameter delta.s is a parameter for insertions.
output	The suffix of output file name. The output contains two files in the working directory. One is the matching nodes and edges between query network and target network, the other is the unique matching node pairs.

#### **Details**

This is an approach for network alignment problem based on conditional random field (CRF) model which uses the node similarity and structure information equally. This method is based on our network querying method net\_query. This method uses an iterative strategy to get the one-to-one map between the query network and target netowrk.

More details can be seen in net\_query.

#### References

Qiang Huang, Ling-Yun Wu, and Xiang-Sun Zhang. CNetA: Network alignment by combining biological and topological features. In Proceedings of 2012 IEEE International Conference on Systems Biology (ISB), 220-225, IEEE, 2012.

Qiang Huang, Ling-Yun Wu, and Xiang-Sun Zhang. Corbi: A new R package for biological network alignment and querying. BMC Systems Biology, 7(Suppl 2):S6, 2013.

#### **Examples**

```
## Not run:
library(Corbi)

## An example: "querynet.txt", "targetnet.txt", "nodesim.txt" are
## three input files in the working directory
net_align("querynet.txt", "targetnet.txt", "nodesim.txt")
## End(Not run)
```

net\_query

Network querying method based on conditional random fields

#### **Description**

Find the best matching subnetworks from a large target network for small query networks based on the conditional random fields (CRF) model.

net\_query

### Usage

```
net_query(
  query.net,
  target.net,
 node.sim,
 query.type = 4,
 delta.d = 1e-10,
  delta.c = 0.5,
  delta.e = 1,
 delta.s = 1,
  output = "result.txt"
)
net_query_batch(
  query.nets,
  target.net,
 node.sim,
 query.type = 4,
  delta.d = 1e-10,
  delta.c = 0.5,
  delta.e = 1,
 delta.s = 1,
  output = "result.txt"
)
```

### Arguments

query.net	The input file name of the query network.
target.net	The input file name of the target network.
node.sim	The input file name of the node similarity scores between the query network and the target network.
query.type	The querying network type: 1 - general, 2 - chain, 3 - tree, 4 - heuristic.
delta.d	The parameter delta.d is a parameter for deletions.
delta.c	The parameter delta.c is a parameter for consecutive deletions.
delta.e	The parameter delta.e is a parameter for single deletion.
delta.s	The parameter delta.s is a parameter for insertions.
output	The suffix of output file name.
query.nets	The vector of input file names of the query networks.

#### **Details**

This is an approach for network querying problem based on conditional random field (CRF) model which can handle both undirected and directed networks, acyclic and cyclic networks, and any number of insertions/deletions.

When querying several networks in the same target network, net\_query\_batch will save much time.

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 query.net: The query network file is written as follows: v1 v2 v3 v4 v5 v3 v4

where v1, v2, v3, v4, v5 ... are the nodes' names and each line indicates there are edges between the first node and other nodes in the line. For example, the first line denotes 4 edges: (v1, v2), (v1, v3), (v1, v4), and (v1, v5).

- target.net: The format of this file is the same as the query network file.
- node.sim: This similarity file's format is as follows: v1 V1 s1

v1 V1 S1 v1 V2 s2

•••

v1 is the node from the query network, V1 is the node from the target network, s1 is the similarity score between the node v1 and V1, and so on.

- query.type: If query.type = 1, the loopy belief propagation (LBP) algorithm will be applied, which is an approximate algorithm for a general graph with loops. If the query is a chain or tree, there are exact algorithms. Set query.type = 2 when the query is a chain, and query.type = 3 when the query is a tree. The heuristic algorithm will be used when query.type = 4, which will try the exact algorithm (junction tree algorithm) first and resort to LBP algorithm when the exact algorithm failed. The default value is 4.
- delta.d: The smaller delta.d is, the heavier penalty for deletions.
- delta.c: The smaller delta.c is, the heavier penalty for consecutive deletions.
- delta.e: The smaller delta.e is, the heavier penalty for single deletion.
- delta.s: The larger delta.s indicates heavier penalty for insertions.

#### References

Qiang Huang, Ling-Yun Wu, and Xiang-Sun Zhang. An Efficient Network Querying Method Based on Conditional Random Fields. Bioinformatics, 27(22):3173-3178, 2011.

### **Examples**

```
## Not run:
library(Corbi)

## An example: "querynet.txt", "targetnet.txt", "nodesim.txt" are
## three input files in the working directory
net_query("querynet.txt", "targetnet.txt", "nodesim.txt", query.type=3)

## End(Not run)

## Not run:
## Batch example
net_query_batch(c("querynet.txt", "querynet2.txt"),
    "targetnet.txt", "nodesim.txt", query.type=3)
```

24 pmultihyper

```
## End(Not run)
```

nnzero

The number of non-zero values of a submatrix

### Description

Retuen the number of non-zero values of the specified submatrix of a given sparse matrix rapidly

### Usage

```
nnzero(m, rows = 1:dim(m)[1], cols = 1:dim(m)[2])
```

### **Arguments**

m	The matrix

rows The integer vector of row index(es) or logical vector indicated the selected rows cols The integer vector of column index(es) or logical vector indicated the selected

cols

### **Details**

This function implements faster calculation algorithm for the CsparseMatrix and RsparseMatrix class in the package **Matrix**.

#### Value

This function will return the number of non-zero values in the specified submatrix.

pmultihyper	The Multivariate Hypergeometric Distribution
pmartingper	The multiured tayper geometric Distribution

### Description

The distribution function for the weighted sums of multivariate hypergeometric distribution

```
pmultihyper(x, k, m, w)
```

pmultinom 25

#### **Arguments**

X	The quantile of weighted sum.
k	The total number of balls drawn from the urn.

m Integer non-negative vector of length N, containing the number of balls of each

color in the urn. N is the number of colors.

w Numeric non-negative vector of length N, specifying the weight of balls of each

color.

#### **Details**

This function gives the distribution function for the weighted sums of multivariate hypergeometric distribution by recursively calling the hypergeometric distribution density function dhyper.

#### Value

This function will return the probablity of  $P(X \le x)$ .

#### See Also

dhyper

pmultinom	The Multinomial Distribution	

### Description

The distribution function for the weighted sums of multinomial distribution

### Usage

```
pmultinom(x, k, m, w)
```

### **Arguments**

x The quantile of weighted sun
--------------------------------

k The total number of balls drawn from the urn.

Mumeric non-negative vector of length N, specifying the probability for drawing the ball of each color; is internally normalized to sum 1. Infinite and missing

values are not allowed. N is the number of colors.

w Numeric non-negative vector of length N, specifying the weight of balls of each

color.

### Details

This function gives the distribution function for the weighted sums of multinomial distribution by recursively calling the binomial distribution density function dbinom.

p\_combine

### Value

This function will return the probablity of  $P(X \le x)$ .

#### See Also

dbinom, dmultinom, rmultinom

|--|

### Description

Combine the statistical significance results from several independent tests by using one of several methods.

### Usage

```
p_combine(p, method = "sumlog", shrink = Inf)
```

### **Arguments**

p the numeric vector containing the p-values need to combine.

method the method use to combine the p-values, can be "sumlog" (Fisher's method),

"sumz" (Stouffer's method).

shrink the number of p-values used in calculation, which are uniform selected from

original p-value vector.

#### Value

This function will return a list with the following components:

p The combined p-value.v The value of statistic.

chisq Use "sumlog" method: The value of chi-squared statistic.

df Use "sumlog" method: The degrees of freedom of chi-squared distribution.

z Use "sumz" method: The value of sum z statistic.

read\_net 27

read\_net

Read network information from text file

### Description

Read the network information from a text file with specific format.

### Usage

```
read_net(file)
```

### **Arguments**

file

The name of text file

### **Details**

This function reads the network information from a text file with specific format: each line contains two strings separated by spaces, which correspond to the names of two end points of one edge in the network.

### Value

A list with the following components:

rode The number of network nodes

The vector of network node names

The logical adjacency matrix

#### See Also

```
write_net
```

rmultihyper

The Multivariate Hypergeometric Distribution

### Description

Generate random variables for the multivariate hypergeometric distribution

```
rmultihyper(n, k, m)
```

28 simulate\_dropout

#### **Arguments**

n	The number	of observations.

k The total number of balls drawn from the urn.

m The integer vector containing the number of balls of each color in the urn.

Length of vector is the number of colors.

#### **Details**

This function generates random variables for the multivariate hypergeometric distribution by iteratively calling hypergeometric random variable generator rhyper.

#### Value

This function will return a matrix of length(m) rows and n columns, and each column contains the number of balls of each color drawn from the urn.

#### See Also

rhyper

simulate\_dropout

Simulate dropout expression data

### Description

Generate the expression data with desired dropout rate

#### Usage

```
simulate_dropout(counts, dropout.rate = 0, dropout.rate.sd = 0.1)
```

#### **Arguments**

counts expression matrix where each row is a gene and each column is a sample.

dropout.rate the desired average dropout rate of all samples.

dropout.rate.sd

the desired standard deviation of dropout rate among samples.

### **Details**

The dropout event is modelled by a logistic distribution such that the low expression genes have higher probability of dropout. The expression value of genes in a sample are randomly set to zero with probabilities associated with their true expression values until the desired dropout rate for that sample is meet.

simulate\_dropout2 29

#### Value

This function will return a list with the following components:

counts The modified expression matrix with the same dimension as input counts.

original.counts

The original input expression matrix.

dropout The binary matrix indicating where the dropout events happen.

#### References

Peter V. Kharchenko, Lev Silberstein, and David T. Scadden. Bayesian approach to single-cell differential expression analysis. Nature Methods, 11(7):740–742, 2014.

simulate\_dropout2

Simulate dropout expression data

### **Description**

Generate the expression data with desired dropout rate range

#### Usage

```
simulate_dropout2(counts, min.rate = 0, max.rate = 0.8)
```

#### **Arguments**

counts expression matrix where each row is a gene and each column is a sample.

min.rate the minimum dropout rate of all samples.

max.rate the maximum dropout rate of all samples.

### **Details**

The dropout event is modelled by a logistic distribution such that the low expression genes have higher probability of dropout. The expression value of genes in a sample are randomly set to zero with probabilities associated with their true expression values until the desired dropout rate for that sample is meet.

#### Value

This function will return a list with the following components:

counts The modified expression matrix with the same dimension as input counts.

original.counts

The original input expression matrix.

dropout The binary matrix indicating where the dropout events happen.

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#### References

Peter V. Kharchenko, Lev Silberstein, and David T. Scadden. Bayesian approach to single-cell differential expression analysis. Nature Methods, 11(7):740–742, 2014.

```
simulate_sample_groups
```

Simulate sample groups from given samples with labels

### **Description**

Generate sample groups with desired labels and sizes from given sample labels.

### Usage

```
simulate_sample_groups(labels, groups, sizes, replace = FALSE)
```

### **Arguments**

labels	a vector containing the label of each sample in the pool.
groups	a vector containing the desired label of samples in each group. The label must be available in the sample pool provided by labels.
sizes	integer vector indicating the desired number of samples in each group. The length must be either one or the same as groups.
replace	logical variable indicating whether sampling is with replacement.

### Value

This function will return a list with the same length as groups. Each component is a vector containing the indexes of samples that are sampled for the corresponding group.

submatrix	Extract a submatrix from a matrix	

### Description

Extract a specified submatrix from a sparse matrix rapidly

```
submatrix(m, rows, cols)
```

URG\_getFactor 31

### **Arguments**

m	The	matrix
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rows The integer vectors of row index(es)

cols The integer vectors of column index(es)

#### **Details**

This function implements faster submatrix extraction algorithm for the CsparseMatrix class in the package **Matrix**.

### Value

This function will return the specified submatrix as a matrix of corresponding type.

URG_getFactor Calculate normalization factors for URG method
URG_getFactor Calculate normalization factors for URG method

#### **Description**

Calculate the normalization factor for each sample by using URG (uniform ratio graph) method.

### Usage

```
URG_getFactor(expr.matrix, p.edge = 0.25, p.gene = 0.4, log.expr = FALSE)
```

### Arguments

expr.matrix	The expression matrix. Each row represents a gene and each column represents a sample.
p.edge	The percentage of gene pairs that are selected into the uniform ratio graph.
p.gene	The maximal percentage of genes that are selected as the stable genes.
log.expr	Logical variable indicating whether the input expression matrix is in logarithmic scale.

#### Value

This function will return a numeric vector with each element [i] represents the normalization factor of sample (i).

### References

Xinhan Ye, Ling-Yun Wu. URG: a new normalization method for gene expression data based on graph model. Manuscript.

### See Also

URG\_normalize

32 write\_net

	- ·	
URG	normalize	

Normalize using given factors

### Description

Normalize the expression matrix by using the given factor for each sample.

### Usage

```
URG_normalize(expr.matrix, factor, log.expr = FALSE)
```

### **Arguments**

expr.matrix The expression matrix. Each row represents a gene and each column represents

a sample.

factor The numeric vector of normalization factors.

log.expr Logical variable indicating whether the input expression matrix is in logarithmic

scale.

#### Value

This function will return a numeric matrix with the same dimension of expr.matrix.

### See Also

URG\_getFactor

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•••	_			

Write network information to text file

### Description

Write the network information to a text file with specific format.

### Usage

```
write_net(net, file)
```

#### **Arguments**

net A list as returned by read\_net

file The name of text file

write\_net 33

### **Details**

This function writes the network information to a text file with specific format: each line contains two strings separated by spaces, which correspond to the names of two end points of one edge in the network.

### See Also

read\_net

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