# Package 'nemTar'

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Type Package
Title A method for singlaing network inference from (epi)genetic aberration profiles
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<b>Description</b> The package 'nemTar' are the extension of classic nest effects models(NEMs)
<b>Depends</b> R ( $\xi$ = 3.5.0), nem ( $\xi$ = 2.60.0), dplyr ( $\xi$ = 0.8.5)
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R topics documented:
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nem\_Tar\_greedy

compute_WIG	Calculating	the	weigheted	information	gain(WIG)	of each	S-
	genes						

## Description

Calculating the weighted information gain(WIG) of each S-genes

## Usage

```
compute_WIG(post_prob, path_affected, num_sgene)
```

## Arguments

post\_prob a list of the probablity that an E-gene is affected by an S-gene with

respect to a specific pathway

path\_affected a list of the E-genes that are affected by the S-genes and are realted to a

specific pathway

num\_sgene a numeric of the number of S-genes

## Value

a list contains the probability of the uniform distribution before the nertwork inference, and the WIG profiles of every S-gene, as well as the WIGs of S-genes

nem_Tar_greedy	Inferring the signaling network from (epi)genetic aberrations and
	effect reporter gene profiles based on greedy hill climbing methods

## Description

Inferring the signaling network from (epi)genetic aberrations and effect reporter gene profiles based on greedy hill climbing methods

## Usage

```
nem_Tar_greedy(D, Sgenes, S_pattern, initial = NULL, control, verbose = TRUE)
```

## Arguments

verbose

D	an E-gene observation matrix with the dimension of p(number of samples)
	$\times$ n(number of effect reporter genes)
Sgenes	a character vector of S-genes
S_pattern	a binary matrix showing the state of whether signaling components (S-genes)are perturbed, with the dimension of m $\times$ m
initial	the initial state of the S-gene graph
control	a list contains necessary parameters for implementing the methods, the details could be referred to ${\sf nem}$

a logic to whether present the execution process

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#### Value

a list contains the network inference results, please also refer to nem

#### See Also

```
nem_Tar_triples nem
```

nem_Tar_triples	Inferring the signaling network from (epi)genetic aberrations and
	effect reporter gene profiles based on triple regition inference

methods

## Description

Inferring the signaling network from (epi)genetic aberrations and effect reporter gene profiles based on triple realtion inference methods

## Usage

```
nem_Tar_triples(D, Sgenes, S_pattern, control, verbose = TRUE)
```

## Arguments

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 $\times$  n(number of effect reporter genes)

Sgenes a character vector of S-genes

S\_pattern a binary matrix showing the state of whether signaling components(S-

genes) are perturbed, with the dimension of m  $\times$  m(m is the number of

S-genes)

control a list contains necessary parameters for implementing the methods, as

nemTar is developed based on nem, the users of nemTar should apply the function nem::set.default.parameters(Sgenes,type="mLL",para=para), para

is a vector of length two:false positive rate and false negative rate.

verbose a logic to whether present the execution process

# Value

a list contains the network inference results using triple relation, please also refer to the nem

#### See Also

```
nem_Tar_greedy nem
```

permu\_test

path_post	To dissect the influenced E-genes and the probability of influence
	under the assumption of nested effects for a specific pathway

# ${\bf Description}$

To dissect the influenced E-genes and the probability of influence under the assumption of nested effects for a specific pathway

# Usage

```
path_post(nemTar_rslt, path_list, Sgenes)
```

# Arguments

nemTar₋rslt	the output of the signaling network inferred by nemTar
$path\_list$	a character of the the signature genes of an pathway
Sgenes	a character of the S-genes within the signaling network

permu_test	Employing the permuation test

# ${\bf Description}$

Employing the permuation test

# ${\bf Usage}$

```
permu_test(path_post, nem_rslt, path_list, sampling_times)
```

# Arguments

path_post	the influenced E-genes and the probability of influence under the assumption of nested effects
nem_rslt	the output of the signaling network inferred by nemTar
$path\_list$	a character of the the signature genes of an pathway
$sampling\_times$	a numeric number indicating the times of sampling

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sampleData2	Generation of the data for sampling in the simulation of the cancer multi-omics samples
	-

## Description

Generation of the data for sampling in the simulation of the cancer multi-omics samples

## Usage

```
sampleData2(
   Phi,
   S_obs,
   p,
   m,
   uninformative = 0,
   type = "binary",
   typeI.err = typeI.err,
   typeII.err = typeII.err)
```

## Arguments

Phi	adjacency matrix
S_obs	binary matrix inddicating the observation of S-genes' states
р	number of samples(patients) to sample
m	number of E-genes to sample
type	an integer, represents the number of row within the raw image
typeI.err	simulated type I error for binary data
typeII.err	simulated type II error for binary data

## See Also

sampleData

 ${\tt WIGsig\_test}$ 

The statistical significance test based on the bootstrap of WIGs

# Description

The statistical significance test based on the bootstrap of WIGs

## Usage

```
WIGsig_test(sample_WIG0, WIG, sampling_times)
```

6 WIG\_sample

#### Arguments

sample\_WIGO the background distribution of the WIG from the sampling of WIG

WIG the pathway-specific WIGcalculated using compute\_WIG sampling\_times a numeric number indicating the times of sampling

#### Value

the statistical significance (BH-adjusted p-value)

WIG\_double Calculating the weighted information gain(WIG) of combinational perturbation of 2 S-genes

## Description

Calculating the weigheted information gain(WIG) of combinational perturbation of 2 S-genes

#### Usage

```
WIG_double(path_post, num_sgene)
```

#### Arguments

path\_post the influenced E-genes and the probability of influence under the assump-

tion of nested effects

num\_sgene a numeric of the number of S-genes

 ${\tt WIG\_sample} \qquad \qquad {\tt Sampling \ of \ the \ WIG \ to \ measure \ the \ statistical \ significance}$ 

# Description

Sampling of the WIG to measure the statistical significance

## Usage

```
WIG_sample(path_post, nem_rslt, path_list, num_sgene, sampling_times)
```

#### Arguments

path\_post the influenced E-genes and the probability of influence under the assump-

tion of nested effects

nem\_rslt the output of the signaling network inferred by nemTar
path\_list a character of the the signature genes of an pathway

num\_sgene a numeric of the number of S-genes

sampling\_times a numeric number indicating the times of sampling

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