# Package 'motif4node'

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Title Motif4node, motif analysis using all non-redundant four-node

Type Package

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<b>Description</b> R package for circuit motif analysis.	
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adj\_to\_tpo

Convert an adjacency matrix to a topology

## Description

Convert an adjacency matrix to a topology

## Usage

```
adj_to_tpo(adj)
```

#### **Arguments**

adj

the adjacency martix to be converted to a topology file

all.circuits

Data file containing the topology of all 60212 non-redundant fournode gene circuits

## Description

This data contains the topology of all 60212 non-redundant four-node gene circuits.

#### Usage

all.circuits

#### **Format**

A list of 60212 circuits, each element is a matrix of adjacency matrix. Four columns/rows represent genes "A", "B", "C", and "D". In each adjacency matrix, 1 represents activation, 2 represents inhibition, and 0 represents no interaction.

analysis\_circuit\_2node

analysis\_circuit\_2node

Analysis script to evaluate the state distribution of a two-node circuit motif

#### Description

Analysis script to evaluate the state distribution of a two-node circuit motif

#### Usage

```
analysis_circuit_2node(tpo, numModels = 10000)
```

#### **Arguments**

tpo Data frame. Topology data of a two-node circuit motif. The data frame contains

three columns: Source, Target, Interaction type

numModels Numeric. Number of models to be simulated. Default: 10000

analysis\_circuit\_4node

Analysis script to evaluate the state distribution of a four-node gene circuit

#### Description

Analysis script to evaluate the state distribution of a four-node gene circuit

#### Usage

```
analysis_circuit_4node(rSet, numModels = 10000)
```

#### **Arguments**

rSet sRACIPE object. RACIPE simulation data.

numModels Numeric. Number of models to be simulated. Default: 10000

#### Value

List. plot objects.

4 dist\_ks

circuit\_grouping

Grouping two-node circuit motifs by their types

#### Description

Grouping two-node circuit motifs by their types

#### Usage

```
circuit_grouping()
```

#### Value

Vector of factor: types for all two-node motifs.

dist\_ks

Calculate th KS distance of two gene expression distributions

#### Description

Calculate th KS distance of two gene expression distributions

#### Usage

```
dist_ks(query, reference, experimental)
```

#### Arguments

query sRACIPE object or PCA matrix of the query data

reference sRACIPE object of the reference data

experimental Logical. T: query is the PCA matrix from an experimental dataset. F: query is

an sRACIPE object

#### Value

the distance between the gene expression distributions from the query and reference.

enrichment\_coupling 5

enrichment_coupling	Motif enrichment analysis for the coupling of two two-node circuit motifs
---------------------	---

## Description

Motif enrichment analysis for the coupling of two two-node circuit motifs

#### Usage

```
enrichment_coupling(
  all.circuits = all.circuits,
  all.scores,
  motif_list,
  new_ind,
  overlap_list,
  decreasing = T,
  if_overlap = 2,
  topCircuits = 600,
  nhill = 20
)
```

#### **Arguments**

all.circuits	List of the topologies of all 60212 non-redundant 4-node circuits. Default "all.circuits" from the package data.
all.scores	Data frame. 1st column: scores; 2nd column: circuit index. The data frame is ordered by the scores.
motif_list	List of 2 by 2 integer matrix. 2-node motif info generated by the function: generate_motif_list.
new_ind	Vector of integer. mapping to non-redundant 2-node motifs generated by the function: generate_index_conversion.
overlap_list	Matrix (6 by 6 integers). Info of overlapping from the motif location in the 4-node circuit generated by the function: generate_overlap_data.
decreasing	Logic. Whether circuits are ranked by the scores in a decreasing order (T) or not (F). Default T.
if_overlap	Whether consider two motifs with overlapping (1), without overlapping (0), or both (2). Default 2.
topCircuits	Integer. Number of top circuits for the enrichment analysis. Default 600.
nhill	Integer/Numeric. Hill coefficient for the Hill function as the weighting factor for motif counts, large n makes the Hill function more binary. Default 20.

#### Value

A data frame containing the enrichment scores of the coupling between 2-node circuit motifs (39 by 39)

```
enrichment_coupling_all_cases
```

A convenient function to perform different motif coupling analyses altogether

#### Description

A convenient function to perform different motif coupling analyses altogether

## Usage

```
enrichment_coupling_all_cases(
  all.circuits,
  all.scores,
  motif_list,
  new_ind,
  overlap_list,
  decreasing = decreasing,
  topCircuits = 600,
  nhill = 20
)
```

## Arguments

all.circuits	List of the topologies of all 60212 non-redundant 4-node circuits. Default "all.circuits from the package data.
all.scores	Data frame. 1st column: scores; 2nd column: circuit index. The data frame is ordered by the scores.
motif_list	List of 2 by 2 integer matrix. 2-node motif info generated by the function: generate_motif_list.
new_ind	Vector of integer. mapping to non-redundant 2-node motifs generated by the function: generate_index_conversion.
overlap_list	Matrix (6 by 6 integers). Info of overlapping from the motif location in the 4-node circuit generated by the function: generate_overlap_data.
decreasing	Logic. Whether circuits are ranked by the scores in a decreasing order (T) or not (F). Default T.
topCircuits	Integer. Number of top circuits for the enrichment analysis. Default 600.
nhill	Integer/Numeric. Hill coefficient for the Hill function as the weighting factor for motif counts, large n makes the Hill function more binary. Default 20.

#### Value

List containing the enrichment of coupling between 2-node motifs for all cases (without overlapping, overlapping, and both)

enrichment\_single 7

enrichment\_single

Motif enrichment analysis for single two-node circuit motifs

#### Description

Motif enrichment analysis for single two-node circuit motifs

#### Usage

```
enrichment_single(
  all.circuits = all.circuits,
  all.scores,
  motif_list,
  new_ind,
  decreasing = T,
  topCircuits = 600,
  nhill = 20
)
```

#### Arguments

all.circuits	List of the topologies of all 60212 non-redundant 4-node circuits. Default "all.circuits" from the package data.
all.scores	Data frame. 1st column: scores; 2nd column: circuit index. The data frame is ordered by the scores.
motif_list	List of 2 by 2 integer matrix. 2-node motif info generated by the function: generate_motif_list.
new_ind	Vector of integer. mapping to non-redundant 2-node motifs generated by the function: generate_index_conversion.
decreasing	Logic. Whether circuits are ranked by the scores in a decreasing order (T) or not (F). Default T.
topCircuits	Integer. Number of top circuits for the enrichment analysis. Default 600.
nhill	Integer/Numeric. Hill coefficient for the Hill function as the weighting factor for motif counts, large n makes the Hill function more binary. Default 20.

#### Value

Data frame containing the enrichment scores of each 2-node circuit motif (39 by 1)

```
{\it Convert\ redundant\ indices\ to\ the\ indices\ for\ all\ 39\ non-redundant\ motifs}
```

#### **Description**

Convert redundant indices to the indices for all 39 non-redundant motifs

#### Usage

```
generate_index_conversion()
```

#### **Description**

Generate a list of all 72 2-node circuit motifs (including redundant ones)

#### Usage

```
generate_motif_list()
```

#### Value

list of all 72 2-node circuit motifs

gen\_network\_scalefree Generate a random scale-free gene network consisting the two-node motifs of choice

#### **Description**

Generate a random scale-free gene network consisting the two-node motifs of choice

#### Usage

```
gen_network_scalefree(num_nodes, motif_list, motif_choice)
```

#### **Arguments**

motif\_list List of 2 by 2 integer matrix. 2-node motif info generated by the function:

generate\_motif\_list.

motif\_choice Vector of integer. A vector of indices of the selected two-node circuit motifs

#### Value

Matrix. Adjacency matrix of the generated network

get\_motif\_adj 9

get_motif_adj  Obtain the adjacency matrix for a specific two-node circuit motion (number should be from 1 to 39)	if
---	----

#### Description

Obtain the adjacency matrix for a specific two-node circuit motif (number should be from 1 to 39)

#### Usage

```
get_motif_adj(number, motif_list)
```

#### Arguments

number Integer. Index of the two-node circuit motif.

motif\_list List of 2 by 2 integer matrix. 2-node motif info generated by the function:

generate\_motif\_list.

#### Value

Integer matrix. 2 by 2 adjacency matrix.

lin_score	The scoring function for ranking circuits with a linear state distribu-
	tion

## Description

The scoring function for ranking circuits with a linear state distribution

## Usage

```
lin_score(rset)
```

#### **Arguments**

rset

The sRACIPE object of the simulated circuit.

#### Value

return(min(score\_vector)): returns the score for the linear state distribution.

10 motif\_analysis

map_and_project	Compare the state distributions of two four-node circuits, find the most matched genes, and project the simulated gene expression data of the 2nd circuit to the PCs of the 1st circuit

#### Description

Compare the state distributions of two four-node circuits, find the most matched genes, and project the simulated gene expression data of the 2nd circuit to the PCs of the 1st circuit

#### Usage

```
map_and_project(rset1, rset2)
```

#### **Arguments**

rset1 sRACIPE object. RACIPE simulation data for the first circuit rset2 sRACIPE object. RACIPE simulation data for the second circuit

#### Value

Numeric matrix. PC coordinates of RACIPE simulated gene expression of the second circuit projected onto the PCs of the RACIPE simulated gene expression of the first circuit

motif\_analysis

A combined motif enrichment analysis for single two-node circuit motifs and motif coupling

## Description

A combined motif enrichment analysis for single two-node circuit motifs and motif coupling

#### Usage

```
motif_analysis(
  all.circuits = all.circuits,
  all.scores,
  ylim = NULL,
  color_breaks = NULL,
  filename = NULL,
  decreasing = T,
  topCircuits = 600
)
```

plot\_adj

#### Arguments

all.circuits	List of the topologies of all $60212$ non-redundant 4-node circuits. Default "all.circuits" from the package data.
all.scores	Data frame. 1st column: scores; 2nd column: circuit index. The data frame is ordered by the scores.
ylim	Vector of numerics (2). Y axis limit for single motif enrichment. Default: NULL.
color_breaks	Vector that defines color scaling for pheatmap. Default: NULL
filename	Character. Prefix of filenames for plotting. Default: NULL. If provided, plots are also saved to files.
decreasing	Logical. Whether circuits are ranked by the scores in a decreasing order (T) or not (F). Default T.
topCircuits	Integer. Number of top circuits for the enrichment analysis. Default 600.

#### Value

List of plotting objects for single motif and motif coupling enrichment analyses.

plot_adj Plot a network from the adjacency matrix	
---	--

## Description

Plot a network from the adjacency matrix

## Usage

```
plot_adj(adj)
```

## Arguments

adj Matrix (4 by 4, integer). Adjacency matrix of a four node circuit.

#### Value

empty

plot\_net

plot\_motif

Plot a specific circuit motif

#### Description

Plot a specific circuit motif

#### Usage

```
plot_motif(number, motif_list)
```

#### **Arguments**

number

Integer. Index of a two-node circuit motif

 $motif\_list$ 

List of 2 by 2 integer matrix. 2-node motif info generated by the function:

generate\_motif\_list.

#### Value

empty

plot\_net

Network plotting function

#### Description

Network plotting function

#### Usage

```
plot_net(tf_links = tf_links)
```

#### Arguments

tf\_links

Data frame of circuit edge list. Three columns: Source, Target, Interaction Types – 1: Activation; 2: Inhibition

#### Value

empty

plot\_RACIPE 13

plot\_RACIPE

Scatterplots of PCA and Gene expression from any 4-node circuit

## Description

Scatterplots of PCA and Gene expression from any 4-node circuit

## Usage

```
plot_RACIPE(rset)
```

#### Arguments

rset

The sRACIPE object of the simulated circuit

#### Value

list(g,p)

simu\_rnorm

RACIPE simulations with random kinetic parameters from Gaussian distributions

## Description

RACIPE simulations with random kinetic parameters from Gaussian distributions

#### Usage

```
simu_rnorm(rset, numModels = 10000)
```

#### Arguments

rset sRACIPE object. sRACIPE output from a standard RACIPE simulation

numModels Numeric. Number of models to be simulated. Default: 10000.

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Simulate one of the 4-node circuits from the 60212 unique circuits

#### Description

Simulate one of the 4-node circuits from the 60212 unique circuits

#### Usage

```
sim_4node(index, Gaussian = F, numModels = 10000, all.circuits = all.circuits)
```

#### **Arguments**

index	Numeric index number of 4-node circuit to be simulate. Takes values from 1:60212
Gaussian	Logical. If T, kinetic parameters will be sampled from a gaussian distribution. If F, kinetic parameters will be sampled from a uniform distribution
numModels	Numeric. Number of models to be simulated. Default: 10000

package data.

#### Value

rset: sRACIPE object. RACIPE simulation results for a circuit

#### **Description**

Generate permutations for p-value calculations

## Usage

```
single_motif_permute(
  all.circuits = all.circuits,
  all.scores,
  decreasing = T,
  topCircuits = 600,
  no_perm
)
```

trig\_score 15

#### **Arguments**

all.circuits	List of the topologies of all 60212 non-redundant 4-node circuits. Default "all.circuits" from the package data.
all.scores	Data frame. 1st column: scores; 2nd column: circuit index. The data frame is ordered by the scores.
decreasing	Logical. Whether circuits are ranked by the scores in a decreasing order (T) or not (F). Default T.
topCircuits	Integer. Number of top circuits for the enrichment analysis. Default 600.
no_perm	Integer. Number of permutations.

#### Value

List of permuted enrichment scores.

trig_score	The scoring function for ranking circuits with a triangular state distribution
------------	--

## Description

The scoring function for ranking circuits with a triangular state distribution

## Usage

```
trig_score(rset)
```

## Arguments

rset The sRACIPE object of the simulated circuit.

#### Value

return(min(score\_vector)): returns the score for the triangular state distribution.

z_score	Calculate z-score for a specific distance	

## Description

Calculate z-score for a specific distance

#### Usage

```
z_score(distances, score)
```

#### Arguments

distances list of distances

score Numeric. score to calculate the z-score

z\_score

## Value

z-score

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