

# Package ‘GeneNetworkBuilder’

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**Type** Package

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**Title** GeneNetworkBuilder: a bioconductor package for building regulatory network using ChIP-chip/ChIP-seq data and Gene Expression Data

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**LinkingTo** Rcpp

**Description** Appliation for discovering direct or indirect targets of transcription factors using ChIP-chip or ChIP-seq, and microarray or RNA-seq gene expression data. Inputting a list of genes of potential targets of one TF from ChIP-chip or ChIP-seq, and the gene expression results, GeneNetworkBuilder generates a regulatory network of the TF.

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GeneNetworkBuilder-package  
*Build Regulatory Network from ChIP-chip/ChIP-seq and Expression Data*

---

**Description**

Appliation for discovering direct or indirect targets of transcription factors using ChIP-chip or ChIP-seq, and microarray or RNA-seq gene expression data. Inputting a list of genes of potential targets of one TF from ChIP-chip or ChIP-seq, and the gene expression results, GeneNetworkBuilder generates a regulatory network of the TF.

**Author(s)**

**Maintainer:** Jianhong Ou Developer <jou@morgridge.org>  
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• Lihua Julie Zhu Developer <Julie.Zhu@umassmed.edu>

---

|               |                       |
|---------------|-----------------------|
| browseNetwork | <i>browse network</i> |
|---------------|-----------------------|

---

## Description

plot network generated by [polishNetwork](#)

## Usage

```
browseNetwork(
  gR = graphNEL(),
  layoutType = c("fdp", "dot", "neato", "twopi", "circo"),
  width = NULL,
  height = NULL,
  maxNodes = 500,
  ...
)
```

## Arguments

|            |                                                                                                   |
|------------|---------------------------------------------------------------------------------------------------|
| gR         | an object of <a href="#">graphNEL</a>                                                             |
| layoutType | layout type. see <a href="#">GraphvizLayouts</a>                                                  |
| width      | width of the figure                                                                               |
| height     | height of the figure                                                                              |
| maxNodes   | max nodes number to plot. Because if there are too many nodes, the running time will be too long. |
| ...        | parameters used by <a href="#">GraphvizLayouts</a>                                                |

## Value

An object of class `htmlwidget` that will intelligently print itself into HTML in a variety of contexts including the R console, within R Markdown documents, and within Shiny output bindings.

## Examples

```
data("ce.miRNA.map")
data("example.data")
data("ce.interactionmap")
data("ce.IDsMap")
sifNetwork<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
cifNetwork<-filterNetwork(rootgene=ce.IDsMap["DAF-16"], sifNetwork=sifNetwork,
  exprsData=uniqueExprsData(example.data$ce.exprData, "Max", condenseName='logFC'),
  mergeBy="symbols",
  miRNAlist=as.character(ce.miRNA.map[, 1]), tolerance=1)
gR<-polishNetwork(cifNetwork)
browseNetwork(gR)
```

---

browseNetwork-shiny      *Shiny bindings for browseNetwork*

---

### Description

Output and render functions for using browseNetwork within Shiny applications and interactive Rmd documents.

### Usage

```
browseNetworkOutput(outputId, width = "100%", height = "400px")

renderBrowseNetwork(expr, env = parent.frame(), quoted = FALSE)
```

### Arguments

|               |                                                                                                                                |
|---------------|--------------------------------------------------------------------------------------------------------------------------------|
| outputId      | output variable to read from                                                                                                   |
| width, height | Must be a valid CSS unit (like '100%', '400px', 'auto') or a number, which will be coerced to a string and have 'px' appended. |
| expr          | An expression that generates a browseNetwork                                                                                   |
| env           | The environment in which to evaluate expr.                                                                                     |
| quoted        | Is expr a quoted expression (with quote())? This is useful if you want to save an expression in a variable.                    |

---

buildNetwork      *construct the regulatory network*

---

### Description

Get all the connections of interesting genes from regulatory map.

### Usage

```
buildNetwork(TFbindingTable, interactionmap, level = 3)
```

### Arguments

|                |                                                                                   |
|----------------|-----------------------------------------------------------------------------------|
| TFbindingTable | a matrix or data.frame with interesting genes. Column names must be 'from', 'to'  |
| interactionmap | Transcription regulatory map. Column names of interactionmap must be 'from', 'to' |
| level          | Depth of node path                                                                |

### Value

a dataframe or matrix of all the connections of interesting genes

**Examples**

```
data("ce.interactionmap")
data("example.data")
xx<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
```

ce.IDsMap

*C.elegans gene name to wormbase identifier map***Description**

map file for converting gene name or sequence name of *Caenorhabditis elegans* to wormbase identifier

**Usage**

```
ce.IDsMap
```

**Format**

character vector

**Details**

character vector with gene name or sequence name as names and wormbase identifier as values.

**Source**

<http://www.wormbase.org/>

**Examples**

```
data(ce.IDsMap)
head(ce.IDsMap)
```

ce.interactionmap

*transcript regulatory map of Caenorhabditis elegans***Description**

transcript regulatory map of *Caenorhabditis elegans*

**Usage**

```
ce.interactionmap
```

**Format**

dataframe

**Details**

transcript regulatory map of *Caenorhabditis elegans* is generated using databases edgedb and microCosm Targets.

**Source**

<http://edgedb.umassmed.edu>, <http://www.ebi.ac.uk/enright-srv/microcosm/htdocs/targets/v5/>

**Examples**

```
data(ce.interactionmap)
head(ce.interactionmap)
```

---

ce.mapIDs

*map file for converting from wormbase identifier to Caenorhabditis elegans gene name*

---

**Description**

map file for converting from wormbase identifier to *Caenorhabditis elegans* gene name

**Usage**

```
ce.mapIDs
```

**Format**

character vector

**Details**

character vector with wormbase identifier as names and gene name as values.

**Source**

<http://www.wormbase.org/>

**Examples**

```
data(ce.mapIDs)
head(ce.mapIDs)
```

---

|              |                                            |
|--------------|--------------------------------------------|
| ce.miRNA.map | <i>micro RNA of Caenorhabditis elegans</i> |
|--------------|--------------------------------------------|

---

**Description**

micro RNA of *Caenorhabditis elegans*

**Usage**

```
ce.miRNA.map
```

**Format**

dataframe

**Details**

The first column is wormbase identifier. And the second column is miRNA names.

**Source**

<http://www.mirbase.org/>

**Examples**

```
data(ce.miRNA.map)
head(ce.miRNA.map)
```

---

|           |                                   |
|-----------|-----------------------------------|
| convertID | <i>convert gene IDs by id map</i> |
|-----------|-----------------------------------|

---

**Description**

For same gene, there are multiple gene alias. In order to eliminate the possibility of missing any connections, convert the gene symbols to unique gene ids is important. This function can convert the gene symbols to unique ids and convert it back according a giving map.

**Usage**

```
convertID(x, IDsMap, ByName = c("from", "to"))
```

**Arguments**

|        |                                                            |
|--------|------------------------------------------------------------|
| x      | a matrix or dataframe contain the columns to be converted. |
| IDsMap | a character vector of the identifier map                   |
| ByName | the column names to be converted                           |

**Value**

a matrix or dataframe with converted gene IDs

**Examples**

```
data("ce.IDsMap")
bind<-cbind(from="daf-16", to=c("fkh-7", "hlh-13", "mxl-3", "nhr-3", "lfi-1"))
convertID(toupper(bind), ce.IDsMap, ByName=c("from", "to"))
```

---

|            |                                     |
|------------|-------------------------------------|
| cy3Network | <i>visualize the network by Cy3</i> |
|------------|-------------------------------------|

---

**Description**

Using RCy3 to visualize the network.

**Usage**

```
cy3Network(
  gR = graphNEL(),
  ...,
  stringify = FALSE,
  species = "Homo sapiens",
  style = "Marquee",
  widths = c(0.25, 5)
)
```

**Arguments**

|           |                                                                                            |
|-----------|--------------------------------------------------------------------------------------------|
| gR        | an object of <a href="#">graphNEL</a>                                                      |
| ...       | parameters will be passed to <a href="#">createNetworkFromGraph</a>                        |
| stringify | Run STRINGify or not                                                                       |
| species   | if stringify is TRUE, teh species will be used to retrieve the nodes and edges properties. |
| style     | The default style when create the network                                                  |
| widths    | The link width range.                                                                      |

**Value**

The network SUID.

## Examples

```
data("ce.miRNA.map")
data("example.data")
data("ce.interactionmap")
data("ce.IDsMap")
sifNetwork<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
cifNetwork<-filterNetwork(rootgene=ce.IDsMap["DAF-16"], sifNetwork=sifNetwork,
  exprsData=uniqueExprsData(example.data$ce.exprData, "Max", condenseName='logFC'),
  mergeBy="symbols",
  miRNAlist=as.character(ce.miRNA.map[, 1]), tolerance=1)
gR<-polishNetwork(cifNetwork)
if(interactive()){
  cy3Network(gR)
}
```

---

example.data

*example datasets for documentation*

---

## Description

example.data is a data list of example datasets. There is a dataset example.microarrayData, which is the example of gene expression data of a gene-chip result of *C.elegans*. Dataset example.data\$ce.bind is a TF binding matrix of ChIP-chip experiment of *C.elegans*. Dataset example.data\$cd.exprData is expression data of a gene-chip result of *C.elegans*. Dataset example.data\$hs.bind is a TF binding matrix of ChIP-chip experiment of *H.sapiens*. Dataset example.data\$hs.exprData is expression data of a combination of a gene-chip result and a RNA-SEQ result of *H.sapiens*.

## Usage

```
example.data
```

## Format

```
dataframe
```

## Details

The dataset example.microarrayData contains columns: ID, logFC, AveExpr, t, P.Value, adj.P.Val, B, genes and symbols. The columns of ID, logFC and symbols are required by GeneNetwork-Builder. The dataset example.data\$hs.bind contains columns: ID, symbols, logFC and P.Value. The dataset example.data\$hs.exprData contains columns: from and to.

## Examples

```
data(example.data)
names(example.data)
head(example.data$example.microarrayData)
head(example.data$ce.bind)
```

```
head(example.data$ce.exprData)
head(example.data$hs.bind)
head(example.data$hs.exprData)
```

---

|               |                                        |
|---------------|----------------------------------------|
| exportNetwork | <i>Save network in various formats</i> |
|---------------|----------------------------------------|

---

## Description

Save graph into HTML, json or xgmml format.

## Usage

```
exportNetwork(network, file, format = c("HTML", "json", "XGMML"), ...)
```

## Arguments

|         |                                                                                                                                             |
|---------|---------------------------------------------------------------------------------------------------------------------------------------------|
| network | output of <a href="#">browseNetwork</a>                                                                                                     |
| file    | Name of the file to save to.                                                                                                                |
| format  | type in which graph shall be saved. Could be one of HTML, json or XGMML.                                                                    |
| ...     | Parameter could be used by <a href="#">saveWidget</a> for HTML or <a href="#">writeLines</a> for json or <a href="#">saveXML</a> for XGMML. |

## Examples

```
data("ce.miRNA.map")
data("example.data")
data("ce.interactionmap")
data("ce.IDsMap")
sifNetwork<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
cifNetwork<-filterNetwork(rootgene=ce.IDsMap["DAF-16"], sifNetwork=sifNetwork,
  exprsData=uniqueExprsData(example.data$ce.exprData, "Max", condenseName='logFC'),
  mergeBy="symbols",
  miRNAlist=as.character(ce.miRNA.map[, 1]), tolerance=1)
gR<-polishNetwork(cifNetwork)
network <- browseNetwork(gR)
exportNetwork(network, "sample.html")
```

---

|               |                                                                  |
|---------------|------------------------------------------------------------------|
| filterNetwork | <i>filter the regulatory network table by expression profile</i> |
|---------------|------------------------------------------------------------------|

---

### Description

verify every nodes in the regulatory network by expression profile

### Usage

```
filterNetwork(
  rootgene,
  sifNetwork,
  exprsData,
  mergeBy = "symbols",
  miRNAlist,
  remove_miRNA = FALSE,
  tolerance = 0,
  cutoffPVal = 0.01,
  cutoffLFC = 0.5,
  minify = TRUE,
  miRNAtol = FALSE
)
```

### Arguments

|              |                                                                                                              |
|--------------|--------------------------------------------------------------------------------------------------------------|
| rootgene     | name of root gene. It must be the ID used in xx regulatory network                                           |
| sifNetwork   | Transcription regulatory network table. Column names of xx must be 'from','to'                               |
| exprsData    | dataset of expression comparison data, which should contain column logFC and column given by exprsDataByName |
| mergeBy      | The column name contains ID information used to merge with 'to' column of sifNetwork in exprsData            |
| miRNAlist    | vector of microRNA ids.                                                                                      |
| remove_miRNA | remove miRNA from the network or not. Bool value, TRUE or FALSE                                              |
| tolerance    | maximum number of unverified nodes in each path                                                              |
| cutoffPVal   | cutoff p value of valid differential expressed gene/miRNA                                                    |
| cutoffLFC    | cutoff log fold change value of a valid differential expressed gene/miRNA                                    |
| minify       | Only keep the best path if multiple paths exists for single node? Bool value, TRUE or FALSE                  |
| miRNAtol     | take miRNA expression into account for tolerance calculation. Bool value, TRUE or FALSE                      |

### Value

a dataframe of filtered regulatory network by expression profile

**Examples**

```

data("ce.miRNA.map")
data("example.data")
data("ce.interactionmap")
data("ce.IDsMap")
sifNetwork<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
cifNetwork<-filterNetwork(rootgene=ce.IDsMap["DAF-16"], sifNetwork=sifNetwork,
  exprsData=uniqueExprsData(example.data$ce.exprData, "Max", condenseName='logFC'),
  mergeBy="symbols",
  miRNAlist=as.character(ce.miRNA.map[, 1]), tolerance=1)

```

---

|           |                                                                                                |
|-----------|------------------------------------------------------------------------------------------------|
| hs.IDsMap | <i>map file for converting gene name or sequence name of Homo sapiens to Entrez identifier</i> |
|-----------|------------------------------------------------------------------------------------------------|

---

**Description**

map file for converting gene name or sequence name of *Homo sapiens* to Entrez identifier

**Usage**

```
hs.IDsMap
```

**Format**

character vector

**Details**

character vector with gene name as names and Entrez identifier as values.

**Examples**

```

data(hs.IDsMap)
head(hs.IDsMap)

```

---

|                   |                                                  |
|-------------------|--------------------------------------------------|
| hs.interactionmap | <i>transcript regulation map of Homo sapiens</i> |
|-------------------|--------------------------------------------------|

---

**Description**

transcript regulation map of *Homo sapiens*

**Usage**

hs.interactionmap

**Format**

dataframe

**Details**

transcript regulatory map of *Homo sapiens* is generated using databases FANTOM, mirGen and microCosm Targets.

**Source**

<http://fantom.gsc.riken.jp/5/>, <http://www.ebi.ac.uk/enright-srv/microcosm/htdocs/targets/v5/>, [http://carolina.imis.athena-innovation.gr/diana\\_tools/web/index.php](http://carolina.imis.athena-innovation.gr/diana_tools/web/index.php)

**Examples**

```
data(hs.interactionmap)
head(hs.interactionmap)
```

---

|           |                                                                                 |
|-----------|---------------------------------------------------------------------------------|
| hs.mapIDs | <i>map file for converting from Entrez identifier to Homo sapiens gene name</i> |
|-----------|---------------------------------------------------------------------------------|

---

**Description**

map file for converting from Entrez identifier to *Homo sapiens* gene name

**Usage**

hs.mapIDs

**Format**

character vector

**Details**

character vector with Entrez identifier as names and gene name as values.

**Examples**

```
data(hs.mapIDs)
head(hs.mapIDs)
```

---

|              |                                  |
|--------------|----------------------------------|
| hs.miRNA.map | <i>micro RNA of Homo sapiens</i> |
|--------------|----------------------------------|

---

**Description**

micro RNA of *Homo sapiens*

**Usage**

```
hs.miRNA.map
```

**Format**

```
dataframe
```

**Details**

The first column is entrez identifier. And the second column is miRNA names.

**Source**

<http://www.mirbase.org/>

**Examples**

```
data(hs.miRNA.map)
head(hs.miRNA.map)
```

---

|                  |                                               |
|------------------|-----------------------------------------------|
| networkFromGenes | <i>Build network by a list of given genes</i> |
|------------------|-----------------------------------------------|

---

### Description

By providing a list of given genes, build a network for input of filterNetwork.

### Usage

```
networkFromGenes(genes, interactionmap, level = 3, unrooted = FALSE)
```

### Arguments

|                |                                                                                  |
|----------------|----------------------------------------------------------------------------------|
| genes          | A vector of character for interested genes.                                      |
| interactionmap | Transcription regulatory map. Column names of interactionmap must be 'from','to' |
| level          | Depth of node path                                                               |
| unrooted       | Return unrooted regulatory network table or not.                                 |

### Value

a list with elements: rootgene: The nodes with maximal connections. sifNetwork: Transcription regulatory network table.

### Examples

```
data("ce.interactionmap")
data("example.data")
genes <- as.character(example.data$ce.bind$from)
xx<-networkFromGenes(example.data$ce.bind, ce.interactionmap, level=2)
```

---

|               |                                                                           |
|---------------|---------------------------------------------------------------------------|
| polishNetwork | <i>generate an object of graphNEL to represent the regulation network</i> |
|---------------|---------------------------------------------------------------------------|

---

### Description

generate an object of graphNEL to represent the regulation network. Each node will has three attributes: size, borderColor and fill. The size will be mapped to the length of its edges. The node fill color will be mapped to logFC.

**Usage**

```
polishNetwork(
  cifNetwork,
  nodeData,
  nodesDefaultSize = 48,
  nodecolor = colorRampPalette(c("green", "yellow", "red"))(5),
  nodeBg = "white",
  nodeBorderColor = list(gene = "darkgreen", miRNA = "darkblue"),
  edgeWeight = NA,
  edgeLwd = 0.5,
  ...
)
```

**Arguments**

|                               |                                                                                                                                        |
|-------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|
| <code>cifNetwork</code>       | dataframe used to draw network graph. column names of <code>cifNetwork</code> must contain 'from', 'to', 'logFC' and 'miRNA'           |
| <code>nodeData</code>         | The node data. If it is not provide, node data will be retrieved from <code>cifNetwork</code> for the 'to' nodes.                      |
| <code>nodesDefaultSize</code> | nodes default size                                                                                                                     |
| <code>nodecolor</code>        | a character vector of color set. The node color will be mapped to color set by log fold change. Or the column names for the colors.    |
| <code>nodeBg</code>           | background of node                                                                                                                     |
| <code>nodeBorderColor</code>  | a list of broder node color set. <code>nodeBorderColor</code> 's element must be gene and miRNA                                        |
| <code>edgeWeight</code>       | the weight of edge. It can be a column name of <code>cifNetwork</code> .                                                               |
| <code>edgeLwd</code>          | the default width of edge. If <code>edgeWeight</code> is set, the <code>edgeLwd</code> will be mapped to the <code>edgeWeight</code> . |
| <code>...</code>              | any parameters can be passed to <a href="#">graph.par</a>                                                                              |

**Value**

An object of `graphNEL` class of the network

**Examples**

```
data("ce.miRNA.map")
data("example.data")
data("ce.interactionmap")
data("ce.IDsMap")
sifNetwork<-buildNetwork(example.data$ce.bind, ce.interactionmap, level=2)
cifNetwork<-filterNetwork(rootgene=ce.IDsMap["DAF-16"], sifNetwork=sifNetwork,
  exprsData=uniqueExprsData(example.data$ce.exprData, "Max", condenseName='logFC'),
  mergeBy="symbols",
  miRNAlist=as.character(ce.miRNA.map[, 1]), tolerance=1)
```

```
gR<-polishNetwork(cifNetwork)
## browseNetwork(gR)
```

---

|           |                              |
|-----------|------------------------------|
| saveXGMML | <i>Save network as xgmml</i> |
|-----------|------------------------------|

---

### Description

Save graph into xgmml format.

### Usage

```
saveXGMML(network, file, ...)
```

### Arguments

|         |                                         |
|---------|-----------------------------------------|
| network | output of <a href="#">browseNetwork</a> |
| file    | Name of the file to save to.            |
| ...     | Parameter could be used by saveXML      |

---

|               |                                  |
|---------------|----------------------------------|
| subsetNetwork | <i>Subset a polished network</i> |
|---------------|----------------------------------|

---

### Description

Subset the output of polishNetwork by a list of nodes name

### Usage

```
subsetNetwork(graph, genes)
```

### Arguments

|       |                                                 |
|-------|-------------------------------------------------|
| graph | A graphNEL object. The output of polishNetwork. |
| genes | A list of nodes names                           |

### Value

An object of graph.

### Examples

```
library(graph)
set.seed(123)
g1 <- randomEGraph(LETTERS[seq.int(15)], edges=100)
g1 <- subsetNetwork(g1, LETTERS[seq.int(5)])
plot(g1)
```

|                 |                                   |
|-----------------|-----------------------------------|
| uniqueExprsData | <i>unique the microarray data</i> |
|-----------------|-----------------------------------|

### Description

get unique the microarray data for each gene id.

## Usage

```
uniqueExprsData(exprsData, method = "Max", condenseName = "logFC")
```

## Arguments

|              |                                       |
|--------------|---------------------------------------|
| exprsData    | dataset of expression comparison data |
| method       | method must be Max, Median or Min     |
| condenseName | column names to be condensed          |

**Value**

a dataframe of expression data without duplicates

## Examples

[illegible]

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