

# Package ‘MSstatsBioNet’

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**Type** Package

**Title** Network Analysis for MS-based Proteomics Experiments

**Version** 1.1.4

**Description** A set of tools for network analysis using mass spectrometry-based proteomics data and network databases. The package takes as input the output of MSstats differential abundance analysis and provides functions to perform enrichment analysis and visualization in the context of prior knowledge from past literature. Notably, this package integrates with INDRA, which is a database of biological networks extracted from the literature using text mining techniques.

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**Depends** R (>= 4.4.0), MSstats

**Imports** RCy3, httr, jsonlite, r2r, tidyR, MASS

**Suggests** data.table, BiocStyle, knitr, rmarkdown, testthat (>= 3.0.0), mockery, MSstatsConvert

**VignetteBuilder** knitr

**biocViews** ImmunoOncology, MassSpectrometry, Proteomics, Software, QualityControl, NetworkEnrichment, Network

**Encoding** UTF-8

**URL** <http://msstats.org>, <https://vitek-lab.github.io/MSstatsBioNet/>

**BugReports** <https://groups.google.com/forum/#!forum/msstats>

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*.populateHgncIdsInDataFrame*  
*Populate HGNC IDs in Data Frame*

---

### Description

This function populates the HGNC IDs in the data frame based on the Uniprot IDs.

### Usage

```
.populateHgncIdsInDataFrame(df)
```

### Arguments

**df** A data frame containing protein information.

### Value

A data frame with populated HGNC IDs.

---

.populateHgncNamesInDataFrame  
*Populate HGNC Names in Data Frame*

---

## Description

This function populates the HGNC names in the data frame based on the HGNC IDs.

## Usage

```
.populateHgncNamesInDataFrame(df)
```

## Arguments

df           A data frame containing protein information.

## Value

A data frame with populated HGNC names.

---

.populateKinaseInfoInDataFrame  
*Populate Kinase Info in Data Frame*

---

## Description

This function populates the kinase information in the data frame based on the HGNC names.

## Usage

```
.populateKinaseInfoInDataFrame(df)
```

## Arguments

df           A data frame containing protein information.

## Value

A data frame with populated kinase information.

---

`.populatePhophataseInfoInDataFrame`  
*Populate Phosphatase Info in Data Frame*

---

## Description

This function populates the phosphatase information in the data frame based on the HGNC names.

## Usage

```
.populatePhophataseInfoInDataFrame(df)
```

## Arguments

`df` A data frame containing protein information.

## Value

A data frame with populated phosphatase information.

---

`.populateTranscriptionFactorInfoInDataFrame`  
*Populate Transcription Factor Info in Data Frame*

---

## Description

This function populates the transcription factor information in the data frame based on the HGNC names.

## Usage

```
.populateTranscriptionFactorInfoInDataFrame(df)
```

## Arguments

`df` A data frame containing protein information.

## Value

A data frame with populated transcription factor information.

---

.populateUniprotIdsInDataFrame  
Populate Uniprot IDs in Data Frame

---

## Description

This function populates the Uniprot IDs in the data frame based on the protein ID type.

## Usage

```
.populateUniprotIdsInDataFrame(df, proteinIdType)
```

## Arguments

df A data frame containing protein information.  
proteinIdType A character string specifying the type of protein ID. It can be either "Uniprot" or "Uniprot\_Mnemonic".

## Value

A data frame with populated Uniprot IDs.

---

.validateAnnotateProteinInfoFromIndraInput  
Validate Annotate Protein Info Input

---

## Description

This function validates the input data frame for the annotateProteinInfoFromIndra function.

## Usage

```
.validateAnnotateProteinInfoFromIndraInput(df)
```

## Arguments

df A data frame containing protein information.

## Value

None. Throws an error if validation fails.

---

**annotateProteinInfoFromIndra**  
*Annotate Protein Information from Indra*

---

## Description

This function annotates a data frame with protein information from Indra.

## Usage

```
annotateProteinInfoFromIndra(df, proteinIdType)
```

## Arguments

<b>df</b>	output of <a href="#">groupComparison</a> function's comparisonResult table, which contains a list of proteins and their corresponding p-values, logFCs, along with additional HGNC ID and HGNC name columns
<b>proteinIdType</b>	A character string specifying the type of protein ID. It can be either "Uniprot" or "Uniprot_Mnemonic".

## Value

A data frame with the following columns:

**Protein** Character. The original protein identifier.

**UniprotID** Character. The Uniprot ID of the protein.

**HgncID** Character. The HGNC ID of the protein.

**HgncName** Character. The HGNC name of the protein.

**IsTranscriptionFactor** Logical. Indicates if the protein is a transcription factor.

**IsKinase** Logical. Indicates if the protein is a kinase.

**IsPhosphatase** Logical. Indicates if the protein is a phosphatase.

## Examples

```
df <- data.frame(Protein = c("CLH1_HUMAN"))
annotated_df <- annotateProteinInfoFromIndra(df, "Uniprot_Mnemonic")
head(annotated_df)
```

---

exportNetworkToHTML     *Export network data with Cytoscape visualization*

---

## Description

Convenience function that takes nodes and edges data directly and creates both the configuration and HTML export in one step.

## Usage

```
exportNetworkToHTML(  
  nodes,  
  edges,  
  filename = "network_visualization.html",  
  displayLabelType = "id",  
  ...  
)
```

## Arguments

nodes	Data frame with node information
edges	Data frame with edge information
filename	Output HTML filename
displayLabelType	Type of label to display ("id" or "hgncName")
...	Additional arguments passed to exportCytoscapeToHTML()

## Value

Invisibly returns the file path of the created HTML file

---

generateCytoscapeConfig  
    *Generate Cytoscape visualization configuration*

---

## Description

This function creates a complete Cytoscape configuration object that can be used to render a network visualization. It's decoupled from any specific UI framework.

## Usage

```
generateCytoscapeConfig(  
  node_elements,  
  edge_elements,  
  container_id = "network-cy",  
  event_handlers = NULL,  
  layout_options = NULL  
)
```

**Arguments**

<code>node_elements</code>	List of node elements created by <code>createNodeElements()</code>
<code>edge_elements</code>	List of edge elements created by <code>createEdgeElements()</code>
<code>container_id</code>	ID of the HTML container element (default: 'network-cy')
<code>event_handlers</code>	Optional list of event handler configurations
<code>layout_options</code>	Optional list of layout configuration options

**Value**

List containing:  
 - elements: Combined node and edge elements  
 - style: Cytoscape style configuration  
 - layout: Layout configuration  
 - container\_id: Container element ID  
 - js\_code: Complete JavaScript code (for backward compatibility)

---

`generateJavaScriptCode`

*Generate JavaScript code from Cytoscape configuration*

---

**Description**

Internal function to convert configuration object to JavaScript code

**Usage**

```
generateJavaScriptCode(config)
```

**Arguments**

<code>config</code>	Configuration object from <code>generateCytoscapeConfig()</code>
---------------------	--

**Value**

Character string containing JavaScript code

---

`getPathwaysFromIndra`    *Get pathways ranked on relevance from INDRA DB***Description**

Get pathways ranked on relevance from INDRA DB

**Usage**

```
getPathwaysFromIndra(
    annotated_df,
    main_target = "MEN1_HUMAN",
    target_type = "Protein"
)
```

**Arguments**

annotated_df	output of <a href="#">groupComparison</a> function's comparisionResult table, which contains a list of proteins and their corresponding p-values, logFCs, along with additional HGNC ID and HGNC name columns
main_target	A main target, e.g. main target of a drug or protein of particular interest
target_type	One of either 'Protein' or 'Drug'. Default is 'Protein'

**Value**

df of pathways

**Examples**

```
annotated_df <- data.table::fread(system.file(  
    "extdata/groupComparisonModel.csv",  
    package = "MSstatsBioNet"  
)  
pathways <- getPathwaysFromIndra(annotated_df, "P05067")  
head(pathways)
```

---

**getSubnetworkFromIndra**

*Get subnetwork from INDRA database*

---

**Description**

Using differential abundance results from MSstats, this function retrieves a subnetwork of protein interactions from INDRA database.

**Usage**

```
getSubnetworkFromIndra(  
    input,  
    protein_level_data = NULL,  
    pvalueCutoff = NULL,  
    statement_types = c("IncreaseAmount", "DecreaseAmount"),  
    paper_count_cutoff = 1,  
    evidence_count_cutoff = 1,  
    correlation_cutoff = 0.3,  
    sources_filter = NULL,  
    logfc_cutoff = NULL,  
    force_include_proteins = NULL,  
    force_include_other = NULL  
)
```

### Arguments

<code>input</code>	output of <code>groupComparison</code> function's <code>comparisionResult</code> table, which contains a list of proteins and their corresponding p-values, logFCs, along with additional HGNC ID and HGNC name columns
<code>protein_level_data</code>	output of the <code>dataProcess</code> function's <code>ProteinLevelData</code> table, which contains a list of proteins and their corresponding abundances. Used for annotating correlation information and applying correlation cutoffs.
<code>pvalueCutoff</code>	p-value cutoff for filtering. Default is NULL, i.e. no filtering
<code>statement_types</code>	list of interaction types to filter on. Equivalent to statement type in INDRA. Default is c("IncreaseAmount", "DecreaseAmount").
<code>paper_count_cutoff</code>	number of papers to filter on. Default is 1.
<code>evidence_count_cutoff</code>	number of evidence to filter on for each paper. E.g. A paper may have 5 sentences describing the same interaction vs 1 sentence. Default is 1.
<code>correlation_cutoff</code>	if <code>protein_level_abundance</code> is not NULL, apply a cutoff for edges with correlation less than a specified cutoff. Default is 0.3
<code>sources_filter</code>	filtering only on specific sources. Default is no filter, i.e. NULL. Otherwise, should be a list, e.g. c('reach', 'medscan').
<code>logfc_cutoff</code>	absolute log fold change cutoff for filtering proteins. Only proteins with  logFC  greater than this value will be retained. Default is NULL, i.e. no logFC filtering.
<code>force_include_proteins</code>	character vector of protein identifiers to exempt from all filtering steps. These proteins will be retained regardless of p-value, logFC, or other filtering criteria. Default is NULL, i.e. no exemptions.
<code>force_include_other</code>	character vector of identifiers to include in the network, regardless if those ids are in the input data. Should be formatted as "namespace:identifier", e.g. "HGNC:1234" or "CHEBI:4911".

### Value

list of 2 data.frames, nodes and edges

### Examples

```
input <- data.table::fread(system.file(
  "extdata/groupComparisonModel.csv",
  package = "MSstatsBioNet"
))
subnetwork <- getSubnetworkFromIndra(input)
head(subnetwork$nodes)
head(subnetwork$edges)
```

---

```
previewNetworkInBrowser
```

*Preview network in browser*

---

## Description

Creates a temporary HTML file and opens it in the default web browser

## Usage

```
previewNetworkInBrowser(nodes, edges, displayLabelType = "id", ...)
```

## Arguments

nodes	Data frame with node information
edges	Data frame with edge information
displayLabelType	Type of label to display ("id" or "hgncName")
...	Additional arguments passed to exportCytoscapeToHTML()

---

```
visualizeNetworks
```

*Create visualization of network*

---

## Description

Use results from INDRA to generate a visualization of the a network on Cytoscape Desktop. Note that the Cytoscape Desktop app must be open for this function to work.

## Usage

```
visualizeNetworks(  
  nodes,  
  edges,  
  pvalueCutoff = 0.05,  
  logfcCutoff = 0.5,  
  node_label_column = "id",  
  main_targets = c()  
)
```

## Arguments

nodes	dataframe of nodes consisting of columns id (character), pvalue (number), logFC (number)
edges	dataframe of edges consisting of columns source (character), target (character), interaction (character), evidenceCount (number), evidenceLink (character)
pvalueCutoff	p-value cutoff for coloring significant proteins. Default is 0.05
logfcCutoff	log fold change cutoff for coloring significant proteins. Default is 0.5

```
node_label_column  
The column of the nodes dataframe to use as the node label. Default is "id".  
"hgncName" can be used for gene name.  
main_targets character vector of main targets to stand-out with a different node shape. Default  
is an empty vector c(). IDs of main targets should match the column used by the  
node_label_column parameter.
```

**Value**

cytoscape visualization of subnetwork

**Examples**

```
input <- data.table::fread(system.file(  
  "extdata/groupComparisonModel.csv",  
  package = "MSstatsBioNet"  
)  
subnetwork <- getSubnetworkFromIndra(input)  
visualizeNetworks(subnetwork$nodes, subnetwork$edges)
```

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