

# Package ‘BioNERO’

April 10, 2023

**Type** Package

**Title** Biological Network Reconstruction Omnibus

**Version** 1.6.1

**Description** BioNERO aims to integrate all aspects of biological network inference in a single package, including data preprocessing, exploratory analyses, network inference, and analyses for biological interpretations. BioNERO can be used to infer gene coexpression networks (GCNs) and gene regulatory networks (GRNs) from gene expression data. Additionally, it can be used to explore topological properties of protein-protein interaction (PPI) networks. GCN inference relies on the popular WGCNA algorithm. GRN inference is based on the “wisdom of the crowds” principle, which consists in inferring GRNs with multiple algorithms (here, CLR, GENIE3 and ARACNE) and calculating the average rank for each interaction pair. As all steps of network analyses are included in this package, BioNERO makes users avoid having to learn the syntaxes of several packages and how to communicate between them. Finally, users can also identify consensus modules across independent expression sets and calculate intra and interspecies module preservation statistics between different networks.

**Depends** R (>= 4.1)

**License** GPL-3

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**LazyData** false

**URL** <https://github.com/almeidasilvaf/BioNERO>

**BugReports** <https://github.com/almeidasilvaf/BioNERO/issues>

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**check\_SFT***Check scale-free topology fit for a given network*

---

**Description**

Check scale-free topology fit for a given network

**Usage**

```
check_SFT(edgelist, net_type = "gcn")
```

**Arguments**

edgelist	Edge list as a data frame containing node 1, node 2 and edge weight.
net_type	Type of biological network. One of "gcn", "grn", or "ppi". Default: gcn.

**Value**

A list with SFT fit statistics and a message indicating if the network is scale-free.

**Examples**

```
set.seed(1)
exp <- t(matrix(rnorm(10000), ncol=1000, nrow=200))
rownames(exp) <- paste0("Gene", 1:nrow(exp))
colnames(exp) <- paste0("Sample", 1:ncol(exp))
cormat <- cor(t(exp))
edges <- cormat_to_edgelist(cormat)
edges <- edges[abs(edges$Weight) > 0.10, ]
check_SFT(edges)
```

**consensus\_modules**      *Identify consensus modules across independent data sets*

**Description**

Identify consensus modules across independent data sets

**Usage**

```
consensus_modules(
  exp_list,
  metadata,
  power,
  cor_method = "spearman",
  net_type = "signed hybrid",
  module_merging_threshold = 0.8,
  verbose = FALSE
)
```

**Arguments**

<b>exp_list</b>	A list containing the expression data frames with genes in row names and samples in column names or ‘SummarizedExperiment’ objects. The list can be created by using <code>list(exp1, exp2, ..., expn)</code> .
<b>metadata</b>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘exp_list’ is a list of ‘SummarizedExperiment’ objects, since the function will extract <code>colData</code> .
<b>power</b>	Numeric vector of beta power for each expression set as calculated by <code>consensus_SFT_fit</code> .
<b>cor_method</b>	Correlation method used for network reconstruction. One of "spearman" (default), "biweight", or "pearson".
<b>net_type</b>	Network type. One of "signed hybrid" (default), "signed" or "unsigned".
<b>module_merging_threshold</b>	Correlation threshold to merge similar modules into a single one. Default: 0.8.
<b>verbose</b>	Logical indicating whether to display progress messages or not. Default: FALSE.

**Value**

A list containing 4 elements:

**consModules** Consensus module assignments  
**consMEs** Consensus module eigengenes  
**exprSize** Description of the multi-set object returned by the function `WGCNA::checkSets`  
**sampleInfo** Metadata for each expression set  
**genes\_cmodules** Data frame of genes and consensus modules  
**dendro\_plot\_objects** Objects to be used in dendrogram plotting

**See Also**

[cutreeDynamic](#)

**Examples**

```
set.seed(12)
data(zma.se)
filt.zma <- filter_by_variance(zma.se, n=500)
zma.set1 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
zma.set2 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
list.sets <- list(zma.set1, zma.set2)
# SFT power previously identified with consensus_SFT_fit()
cons_mod <- consensus_modules(list.sets, power = c(11, 13),
                                cor_method = "pearson")
```

consensus\_SFT\_fit      *Pick power to fit networks to scale-free topology*

**Description**

Pick power to fit networks to scale-free topology

**Usage**

```
consensus_SFT_fit(
  exp_list,
  setLabels = NULL,
  metadata = NULL,
  cor_method = "spearman",
  net_type = "signed hybrid",
  rsquared = 0.8
)
```

## Arguments

<code>exp_list</code>	A list of expression data frames or SummarizedExperiment objects. If input is a list of data frames, row names must correspond to gene IDs and column names to samples. The list can be created with <code>list(exp1, exp2, ..., expn)</code> .
<code>setLabels</code>	Character vector containing labels for each expression set.
<code>metadata</code>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘ <code>exp_list</code> ’ is a list of ‘SummarizedExperiment’ objects, since the function will extract <code>colData</code> .
<code>cor_method</code>	Correlation method used for network reconstruction. One of "spearman" (default), "biweight", or "pearson".
<code>net_type</code>	Network type. One of "signed hybrid" (default), "signed" or "unsigned".
<code>rsquared</code>	Minimum R squared to consider the network similar to a scale-free topology. Default is 0.8.

## Value

A list of 2 elements:

**power** Numeric vector of optimal beta powers to fit networks to SFT

**plot** A ggplot object displaying main statistics of the SFT fit test

## Examples

```
set.seed(12)
data(zma.se)
filt.zma <- filter_by_variance(zma.se, n=500)
zma.set1 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
zma.set2 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
list.sets <- list(zma.set1, zma.set2)
cons_sft <- consensus_SFT_fit(list.sets, setLabels = c("Maize1", "Maize2"),
                               cor_method = "pearson")
```

<code>consensus_trait_cor</code>	<i>Correlate set-specific modules and consensus modules to sample information</i>
----------------------------------	---

## Description

Correlate set-specific modules and consensus modules to sample information

## Usage

```
consensus_trait_cor(
  consensus,
  cor_method = "spearman",
  continuous_trait = FALSE,
  palette = "RdYlBu",
  cex.lab.x = 0.6,
  cex.lab.y = 0.6,
  cex.text = 0.6,
  transpose = FALSE
)
```

## Arguments

consensus	Consensus network returned by <code>consensus_modules</code> .
cor_method	Correlation method to be used. One of 'spearman' or 'pearson'. Default is 'spearman'.
continuous_trait	Logical indicating if trait is a continuous variable. Default is FALSE.
palette	RColorBrewer's color palette to use. Default is "RdYlBu", a palette ranging from blue to red.
cex.lab.x	Font size for x axis labels. Default: 0.6.
cex.lab.y	Font size for y axis labels. Default: 0.6.
cex.text	Font size for numbers inside matrix. Default: 0.6.
transpose	Logical indicating whether to transpose the heatmap of not. Default is FALSE.

## Details

Significance levels: 1 asterisk: significant at alpha = 0.05. 2 asterisks: significant at alpha = 0.01. 3 asterisks: significant at alpha = 0.001. no asterisk: not significant.

## Value

Data frame of consensus module-trait correlations and p-values.

## See Also

[corPValueFisher](#), [labels2colors](#), [labeledHeatmap](#), [blueWhiteRed](#)

## Examples

```
set.seed(12)
data(zma.se)
filt.zma <- filter_by_variance(zma.se, n=500)
zma.set1 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
zma.set2 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
list.sets <- list(zma.set1, zma.set2)
```

```
# SFT power previously identified with consensus_SFT_fit()
consensus <- consensus_modules(list.sets, power = c(11, 13),
                               cor_method = "pearson")
consensus_trait <- consensus_trait_cor(consensus, cor_method = "pearson")
```

cormat\_to\_edgelist      *Transform a correlation matrix to an edge list*

### Description

Transform a correlation matrix to an edge list

### Usage

```
cormat_to_edgelist(matrix)
```

### Arguments

matrix	Symmetrical correlation matrix.
--------	---------------------------------

### Value

A 2-column data frame containing node 1, node 2 and edge weight.

### Examples

```
data(filt.se)
cor_mat <- cor(t(SummarizedExperiment::assay(filt.se)))
edgelist <- cormat_to_edgelist(cor_mat)
```

detect\_communities      *Detect communities in a network*

### Description

Detect communities in a network

### Usage

```
detect_communities(edgelist, method = igraph::cluster_infomap, directed = TRUE)
```

## Arguments

<code>edgelist</code>	Data frame containing the network as an edge list. First column must be node 1 and second column must be node 2. Additional columns will be interpreted as edge attributes and will be modified by this function.
<code>method</code>	<code>igraph</code> function to be used for community detection. Available functions are <code>cluster_infomap</code> , <code>cluster_edge_betweenness</code> , <code>cluster_fast_greedy</code> , <code>cluster_walktrap</code> , <code>cluster_springlass</code> , <code>cluster_leading_eigen</code> , <code>cluster_louvain</code> , and <code>cluster_label_prop</code> . Default is <code>cluster_infomap</code> .
<code>directed</code>	Logical indicating whether the network is directed (GRN only) or not (GCN and PPI networks). Default: TRUE.

## Value

A data frame containing node names in the first column, and communities to which nodes belong in the second column.

## Author(s)

Fabricio Almeida-Silva

## See Also

[cluster\\_infomap](#), [cluster\\_edge\\_betweenness](#), [cluster\\_fast\\_greedy](#), [cluster\\_walktrap](#), [cluster\\_springlass](#), [cluster\\_leading\\_eigen](#), [cluster\\_louvain](#), [cluster\\_label\\_prop](#)

## Examples

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_edges <- grn_infer(filt.se, method = "clr", regulators = tfs)
com <- detect_communities(grn_edges, directed=TRUE)
```

dfs2zone

*Combine multiple expression tables (.tsv) into a single data frame*

## Description

This function reads multiple expression tables (.tsv files) in a directory and combines them into a single gene expression data frame.

## Usage

```
dfs2zone(mypath, pattern = ".tsv$")
```

## Arguments

<code>mypath</code>	Path to directory containing .tsv files. Files must have the first column in common, e.g. "Gene_ID". Rows are gene IDs and columns are sample names.
<code>pattern</code>	Pattern contained in each expression file. Default is '.tsv\$', which means that all files ending in '.tsv' in the specified directory will be considered expression files.

## Value

Data frame with gene IDs as row names and their expression values in each sample (columns).

## Author(s)

Fabricio Almeida-Silva

## Examples

```
# Simulate two expression data frames of 100 genes and 30 samples
genes <- paste0(rep("Gene", 100), 1:100)
samples1 <- paste0(rep("Sample", 30), 1:30)
samples2 <- paste0(rep("Sample", 30), 31:60)
exp1 <- cbind(genes, as.data.frame(matrix(rnorm(100*30), nrow=100, ncol=30)))
exp2 <- cbind(genes, as.data.frame(matrix(rnorm(100*30), nrow=100, ncol=30)))
colnames(exp1) <- c("Gene", samples1)
colnames(exp2) <- c("Gene", samples2)

# Write data frames to temporary files
tmpdir <- tempdir()
tmp1 <- tempfile(tmpdir = tmpdir, fileext = ".exp.tsv")
tmp2 <- tempfile(tmpdir = tmpdir, fileext = ".exp.tsv")
write.table(exp1, file=tmp1, quote=FALSE, sep="\t")
write.table(exp2, file=tmp2, quote=FALSE, sep="\t")

# Load the files into one
exp <- dfsZone(mypath = tmpdir, pattern=".exp.tsv")
```

## Description

Perform enrichment analysis for a set of genes

**Usage**

```
enrichment_analysis(
  genes,
  background_genes,
  annotation,
  column = NULL,
  correction = "BH",
  p = 0.05,
  bp_param = BiocParallel::SerialParam()
)
```

**Arguments**

<code>genes</code>	Character vector containing genes for overrepresentation analysis.
<code>background_genes</code>	Character vector of genes to be used as background for the Fisher's Exact Test.
<code>annotation</code>	Annotation data frame with genes in the first column and functional annotation in the other columns. This data frame can be exported from Biomart or similar databases.
<code>column</code>	Column or columns of annotation to be used for enrichment. Both character or numeric values with column indices can be used. If users want to supply more than one column, input a character or numeric vector. Default: all columns from <code>annotation</code> .
<code>correction</code>	Multiple testing correction method. One of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr" or "none". Default is "BH".
<code>p</code>	P-value threshold. P-values below this threshold will be considered significant. Default is 0.05.
<code>bp_param</code>	BiocParallel back-end to be used. Default: <code>BiocParallel::SerialParam()</code>

**Value**

Data frame containing significant terms, p-values and associated genes.

**Author(s)**

Fabricio Almeida-Silva

**Examples**

```
data(filt.se)
data(zma.interpro)
genes <- rownames(filt.se)[1:50]
background_genes <- rownames(filt.se)
annotation <- zma.interpro
# Using p = 1 to show all results
enrich <- enrichment_analysis(genes, background_genes, annotation, p = 1)
```

---

exp2gcn*Reconstruct gene coexpression network from gene expression*

---

## Description

Reconstruct gene coexpression network from gene expression

## Usage

```
exp2gcn(
  exp,
  net_type = "signed",
  module_merging_threshold = 0.8,
  SFTpower = NULL,
  cor_method = "spearman",
  verbose = FALSE
)
```

## Arguments

exp	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
net_type	Network type. One of 'signed', 'signed hybrid' or 'unsigned'. Default: 'signed'.
module_merging_threshold	Correlation threshold to merge similar modules into a single one. Default: 0.8.
SFTpower	SFT power generated by the function <code>SFT_fit</code> .
cor_method	Correlation method. One of "pearson", "biweight" or "spearman". Default is "spearman".
verbose	Logical indicating whether to display progress messages or not. Default: FALSE.

## Value

List containing:

- Adjacency matrix
- Data frame of module eigengenes
- Data frame of genes and their corresponding modules
- Data frame of intramodular connectivity
- Correlation matrix
- Parameters used for network reconstruction
- Objects to plot the dendrogram in `plot_dendro_and_colors`.

## Author(s)

Fabricio Almeida-Silva

**See Also**

[adjacency](#), [fromSimilarity](#), [TOMsimilarity](#), [standardColors](#), [labels2colors](#), [moduleEigengenes](#), [plotEigengeneNetwork](#), [cutreeDynamicTree](#)

**Examples**

```
data(filt.se)
# The SFT fit was previously calculated and the optimal power was 16
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
```

exp2grn

*Infer gene regulatory network from expression data***Description**

Infer gene regulatory network from expression data

**Usage**

```
exp2grn(
  exp,
  regulators = NULL,
  eps = 0,
  estimator_aracne = "spearman",
  estimator_clr = "pearson",
  remove_zero = TRUE,
  nsplit = 10,
  ...
)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>regulators</code>	A character vector of regulators (e.g., transcription factors or miRNAs). All regulators must be included in ‘exp’.
<code>eps</code>	Numeric value indicating the threshold used when removing an edge: for each triplet of nodes (i,j,k), the weakest edge, say (ij), is removed if its weight is below $\min(ij, jk) - \text{eps}$ . Default: 0.
<code>estimator_aracne</code>	Entropy estimator to be used in ARACNE inference. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "spearman".
<code>estimator_clr</code>	Entropy estimator to be used in CLR inference. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "pearson".

remove_zero	Logical indicating whether to remove edges whose weight is exactly zero. Zero values indicate edges that were removed by ARACNE. Default: TRUE.
nsplit	Number of groups in which the edge list will be split. Default: 10.
...	Additional arguments passed to ‘GENIE3::GENIE3()’.

## Details

This function infers GRNs with ARACNE, GENIE3 and CLR, ranks correlation weights for each GRN and calculates the average rank for each edge. Then, the resulting GRN is filtered to keep the top n edges that lead to the optimal scale-free topology fit.

## Value

A filtered edge list with regulators in the first column and targets in the second column.

## Examples

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
# Test with small number of trees for demonstration purpose
grn <- exp2grn(filt.se, regulators = tfs, nTrees=2, nsplit=2)
```

**exp\_genes2orthogroups** *Collapse gene-level expression data to orthogroup level*

## Description

For a given list of expression data, this function replaces genes with their corresponding orthogroups to allow inter-species comparisons.

## Usage

```
exp_genes2orthogroups(expList = NULL, og = NULL, summarize = "median")
```

## Arguments

expList	List of expression data frames or SummarizedExperiment objects.
og	Data frame of 3 columns corresponding to orthogroup, species ID, and gene ID, respectively. Species IDs must be the same as the names of the expression list.
summarize	Centrality measure to summarize multiple paralogous genes in the same orthogroup. One of "median" or "mean". Default: "median".

## Value

List of expression data frames for each species with expression summarized at the orthogroup level.

## Examples

```
data(og.zma.osa)
data(zma.se)
data(osa.se)
expList <- list(zma = zma.se,
                 osa = osa.se)
og <- og.zma.osa
exp_ortho <- exp_genes2orthogroups(expList, og, summarize = "mean")
```

exp\_preprocess

*Preprocess expression data for network reconstruction*

## Description

Preprocess expression data for network reconstruction

## Usage

```
exp_preprocess(
  exp,
  NA_rm = TRUE,
  replaceby = 0,
  Zk_filtering = TRUE,
  zk = -2,
  cor_method = "spearman",
  remove_nonexpressed = TRUE,
  method = "median",
  min_exp = 1,
  min_percentage_samples = 0.25,
  remove_confounders = TRUE,
  variance_filter = FALSE,
  n = NULL,
  percentile = NULL,
  vtransform = FALSE
)
```

## Arguments

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a 'SummarizedExperiment' object.
<code>NA_rm</code>	Logical. It specifies whether to remove missing values from the expression data frame or not. Default = TRUE.
<code>replaceby</code>	If <code>NA_rm</code> is TRUE, what to use instead of NAs. One of 0 or 'mean'. Default is 0.
<code>Zk_filtering</code>	Logical. It specifies whether to filter outlying samples by Zk or not. Default: TRUE.

<b>zk</b>	If Zk_filtering is TRUE, the standardized connectivity threshold. Samples below this threshold will be considered outliers. Default is -2.
<b>cor_method</b>	If Zk_filtering is TRUE, the correlation method to use. One of 'spearman', 'bicor', or 'pearson'. Default is 'spearman'.
<b>remove_nonexpressed</b>	Logical. It specifies whether non-expressed genes should be removed or not. Default is TRUE.
<b>method</b>	If remove_nonexpressed is TRUE, the criterion to filter non-expressed genes out. One of "mean", "median", "percentage", or "allsamples". Default is 'median'.
<b>min_exp</b>	If method is 'mean', 'median', or 'allsamples', the minimum value for a gene to be considered expressed. If method is 'percentage', the minimum value each gene must have in at least n percent of samples to be considered expressed.
<b>min_percentage_samples</b>	If method is 'percentage', expressed genes must have expression $\geq$ min_exp in at least this percentage. Values must range from 0 to 1. Default = 0.25.
<b>remove_cofounders</b>	Logical. If TRUE, it removes principal components that add noise to the data.
<b>variance_filter</b>	Logical. If TRUE, it will filter genes by variance. Default is FALSE.
<b>n</b>	If variance_filter is TRUE, the number of most variable genes to keep.
<b>percentile</b>	If variance_filter is TRUE, the percentage of most variable genes to keep.
<b>vstransform</b>	Logical indicating if data should be variance stabilizing transformed. This parameter can only be set to TRUE if data is a matrix of raw read counts.

## Value

Processed gene expression data frame with gene IDs in row names and sample names in column names or 'SummarizedExperiment' object.

## Author(s)

Fabricio Almeida-Silva

## References

Love, M. I., Huber, W., & Anders, S. (2014). Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome biology*, 15(12), 1-21.

## See Also

[varianceStabilizingTransformation](#)

## Examples

```
data(zma.se)
exp <- exp_preprocess(zma.se, variance_filter=TRUE, n=1000)
```

---

**filt.se***Filtered maize gene expression data from Shin et al., 2021.*

---

**Description**

Filtered expression data in transcripts per million (TPM) from Shin et al., 2021. This is the same data set described in zma.se, but it only contains the top 500 genes with the highest variances. This data set was created to be used in unit tests and examples.

**Usage**

```
data(filt.se)
```

**Format**

An object of class `SummarizedExperiment`

**References**

Shin, J., Marx, H., Richards, A., Vaneechoutte, D., Jayaraman, D., Maeda, J., ... & Roy, S. (2021). A network-based comparative framework to study conservation and divergence of proteomes in plant phylogenies. *Nucleic Acids Research*, 49(1), e3-e3.

**Examples**

```
data(filt.se)
```

---

**filter\_by\_variance***Keep only genes with the highest variances*

---

**Description**

Keep only genes with the highest variances

**Usage**

```
filter_by_variance(exp, n = NULL, percentile = NULL)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>n</code>	Number of most variable genes (e.g., <code>n=5000</code> will keep the top 5000 most variable genes).
<code>percentile</code>	Percentile of most highly variable genes (e.g., <code>percentile=0.1</code> will keep the top 10 percent most variable genes). Values must range from 0 to 1.

**Value**

Expression data frame or ‘SummarizedExperiment‘ object with the most variable genes in row names and samples in column names.

**Author(s)**

Fabricio Almeida-Silva

**Examples**

```
data(zma.se)
filt_exp <- filter_by_variance(zma.se, p=0.1)
```

<i>gene_significance</i>	<i>Calculate gene significance for a given group of genes</i>
--------------------------	---

**Description**

Calculate gene significance for a given group of genes

**Usage**

```
gene_significance(
  exp,
  metadata,
  genes = NULL,
  alpha = 0.05,
  min_cor = 0.2,
  use_abs = TRUE,
  palette = "RdYlBu",
  show_rownames = FALSE,
  continuous_trait = FALSE
)
```

**Arguments**

<b>exp</b>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment‘ object.
<b>metadata</b>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘exp‘ is a ‘SummarizedExperiment‘ object, since the function will extract colData.
<b>genes</b>	Character vector of genes to be correlated with traits. If not given, all genes in ‘exp‘ will be considered.
<b>alpha</b>	Significance level. Default is 0.05.
<b>min_cor</b>	Minimum correlation coefficient. Default is 0.2.

use_abs	Logical indicating whether to filter by correlation using absolute value or not. If TRUE, a min_cor of say 0.2 would keep all correlations above 0.2 and below -0.2. Default is TRUE.
palette	RColorBrewer's color palette to use. Default is "RdYlBu", a palette ranging from blue to red.
show_rownames	Logical indicating whether to show row names or not. Default is FALSE.
continuous_trait	Logical indicating if trait is a continuous variable. Default is FALSE.

**Value**

A heatmap of gene significance (GS) and a list containing:

- filtered\_corandpFiltered matrix of correlation and p-values
- raw\_GSRaw matrix of gene significances

**See Also**

[corPValueStudent](#) [RColorBrewer](#)

**Examples**

```
data(filt.se)
gs <- gene_significance(filt.se)
```

get_edge_list	<i>Get edge list from an adjacency matrix for a group of genes</i>
---------------	--

**Description**

Get edge list from an adjacency matrix for a group of genes

**Usage**

```
get_edge_list(
  net,
  genes = NULL,
  module = NULL,
  filter = FALSE,
  method = "optimalsFT",
  r_optimal_test = seq(0.4, 0.9, by = 0.1),
  Zcutoff = 1.96,
  pvalue_cutoff = 0.05,
  rcutoff = 0.7,
  nSamples = NULL,
  check_SFT = FALSE,
  bp_param = BiocParallel::SerialParam()
)
```

## Arguments

<code>net</code>	List object returned by <code>exp2gcn</code> .
<code>genes</code>	Character vector containing a subset of genes from which edges will be extracted. It can be ignored if the user wants to extract an edge list for a given module instead of individual genes.
<code>module</code>	Character with module name from which edges will be extracted. To include 2 or more modules, input the names in a character vector.
<code>filter</code>	Logical indicating whether to filter the edge list or not.
<code>method</code>	Method to filter spurious correlations. One of "Zscore", "optimalSFT", "pvalue" or "min_cor". See details for more information on the methods. Default: 'optimalSFT'
<code>r_optimal_test</code>	Numeric vector with the correlation thresholds to be tested for optimal scale-free topology fit. Only valid if <code>method</code> equals "optimalSFT". Default: <code>seq(0.4, 0.9, by = 0.1)</code>
<code>Zcutoff</code>	Minimum Z-score threshold. Only valid if <code>method</code> equals "Zscore". Default: 1.96
<code>pvalue_cutoff</code>	Maximum P-value threshold. Only valid if <code>method</code> equals "pvalue". Default: 0.05
<code>rcutoff</code>	Minimum correlation threshold. Only valid if <code>method</code> equals "min_cor". Default: 0.7
<code>nSamples</code>	Number of samples in the data set from which the correlation matrix was calculated. Only required if <code>method</code> equals "pvalue".
<code>check_SFT</code>	Logical indicating whether to test if the resulting network is close to a scale-free topology or not. Default: FALSE.
<code>bp_param</code>	BiocParallel back-end to be used. Default: <code>BiocParallel::SerialParam()</code>

## Details

The default method ("optimalSFT") will create several different edge lists by filtering the original correlation matrix by the thresholds specified in `r_optimal_test`. Then, it will calculate a scale-free topology fit index for each of the possible networks and return the network that best fits the scale-free topology. The method "Zscore" will apply a Fisher Z-transformation for the correlation coefficients and remove the Z-scores below the threshold specified in `Zcutoff`. The method "pvalue" will calculate Student asymptotic p-value for the correlations and remove correlations whose p-values are above the threshold specified in `pvalue_cutoff`. The method "min\_cor" will remove correlations below the minimum correlation threshold specified in `rcutoff`.

## Value

Data frame with edge list for the input genes.

## Author(s)

Fabricio Almeida-Silva

**See Also**

[scaleFreeFitIndex](#),[corPValueStudent](#) [fit\\_power\\_law](#)  
[SFT\\_fit](#)  
[exp2gcn.](#)

**Examples**

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
genes <- rownames(filt.se)[1:50]
edges <- get_edge_list(gcn, genes=genes, filter = FALSE)
```

---

get\_HK*Get housekeeping genes from global expression profile*

---

**Description**

Get housekeeping genes from global expression profile

**Usage**

```
get_HK(exp)
```

**Arguments**

**exp** A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.

**Details**

This function identifies housekeeping genes, which are broadly expressed genes with low variation in a global scale across samples. For some cases, users would want to remove these genes as they are not interesting for coexpression network analyses. See references for more details.

**Value**

Character vector of housekeeping gene IDs.

**Author(s)**

Fabricio Almeida-Silva

**References**

Machado, F.B., Moharana, K.C., Almeida-Silva, F., Gazara, R.K., Pedrosa-Silva, F., Coelho, F.S., Grativil, C. and Venancio, T.M. (2020), Systematic analysis of 1298 RNA-Seq samples and construction of a comprehensive soybean (*Glycine max*) expression atlas. *Plant J.* 103: 1894-1909.

**Examples**

```
data(zma.se)
hk <- get_HK(zma.se)
```

**get\_hubs\_gcn**

*Get GCN hubs*

**Description**

Get GCN hubs

**Usage**

```
get_hubs_gcn(exp, net)
```

**Arguments**

- |                  |  |
|------------------|--|
| <code>exp</code> | A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object. |
| <code>net</code> | List object returned by <code>exp2gcn</code> .   |

**Value**

Data frame containing gene IDs, modules and intramodular connectivity of all hubs.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[signedKME](#)

**Examples**

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
hubs <- get_hubs_gcn(filt.se, gcn)
```

---

get_hubs_grn	<i>Get hubs for gene regulatory network</i>
--------------	---

---

## Description

Get hubs for gene regulatory network  
Get hubs for protein-protein interaction network

## Usage

```
get_hubs_grn(  
  edgelist,  
  top_percentile = 0.1,  
  top_n = NULL,  
  return_degree = FALSE,  
  ranked = TRUE  
)  
  
get_hubs_ppi(  
  edgelist,  
  top_percentile = 0.1,  
  top_n = NULL,  
  return_degree = FALSE  
)
```

## Arguments

edgelist	A protein-protein interaction network represented as an edge list.
top_percentile	Numeric from 0 to 1 indicating the percentage of proteins with the highest degree to consider hubs. Default: 0.1.
top_n	Numeric indicating the number of proteins with the highest degree to consider hubs.
return_degree	Logical indicating whether to return a data frame of degree for all proteins. If TRUE, the function will return a list instead of a data frame. Default: FALSE.
ranked	Logical indicating whether to treat third column of the edge list (edge weights) as ranked values. Ignored if the edge list only contains 2 columns. Default: TRUE.

## Value

A data frame with gene ID in the first column and out degree in the second column or a list of two data frames with hubs and degree for all genes, respectively.

A data frame with protein ID in the first column and degree in the second column or a list of two data frames with hubs and degree for all genes, respectively.

## Examples

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_list <- grn_combined(filt.se, regulators=tfs, nTrees=2)
ranked_grn <- grn_average_rank(grn_list)
# split in only 2 groups for demonstration purposes
filtered_edges <- grn_filter(ranked_grn, nsplit=2)
hubs <- get_hubs_grn(filtered_edges)
ppi_edges <- igraph::get.edgelist(igraph::barabasi.game(n=500, directed=FALSE))
hubs <- get_hubs_ppi(ppi_edges, return_degree = TRUE)
```

**get\_neighbors**

*Get 1st-order neighbors of a given gene or group of genes*

## Description

Get 1st-order neighbors of a given gene or group of genes

## Usage

```
get_neighbors(genes, net, cor_threshold = 0.7)
```

## Arguments

<b>genes</b>	Character vector containing genes from which direct neighbors will be extracted.
<b>net</b>	List object returned by exp2gcn.
<b>cor_threshold</b>	Correlation threshold to filter connections. As a weighted network is a fully connected graph, a cutoff must be selected. Default is 0.7.

## Value

List containing 1st-order neighbors for each input gene.

## Author(s)

Fabricio Almeida-Silva

## See Also

`exp2gcn` `SFT_fit`

## Examples

```
data(filt.se)
genes <- rownames(filt.se)[1:10]
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
neighbors <- get_neighbors(genes, gcn)
```

---

grn_average_rank	<i>Rank edge weights for GRNs and calculate average across different methods</i>
------------------	--

---

**Description**

Rank edge weights for GRNs and calculate average across different methods

**Usage**

```
grn_average_rank(list_edges)
```

**Arguments**

list\_edges      List containing edge lists as returned by the function `grn_combined`.

**Value**

Edge list containing regulator, target and mean rank from all algorithms.

**Examples**

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_list <- grn_combined(filt.se, regulators=tfs, nTrees=2)
ranked_grn <- grn_average_rank(grn_list)
```

---

grn_combined	<i>Infer gene regulatory network with multiple algorithms and combine results in a list</i>
--------------	---

---

**Description**

Infer gene regulatory network with multiple algorithms and combine results in a list

**Usage**

```
grn_combined(
  exp,
  regulators = NULL,
  eps = 0.1,
  estimator_aracne = "spearman",
  estimator_clr = "pearson",
  remove_zero = TRUE,
  ...
)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a 'SummarizedExperiment' object.
<code>regulators</code>	A character vector of regulators (e.g., transcription factors or miRNAs). All regulators must be included in 'exp'.
<code>eps</code>	Numeric value indicating the threshold used when removing an edge: for each triplet of nodes (i,j,k), the weakest edge, say (ij), is removed if its weight is below $\min(ij, jk) - \text{eps}$ . Default: 0.1.
<code>estimator_aracne</code>	Entropy estimator to be used in ARACNE inference. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "spearman".
<code>estimator_clr</code>	Entropy estimator to be used in CLR inference. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "pearson".
<code>remove_zero</code>	Logical indicating whether to remove edges whose weight is exactly zero. Zero values indicate edges that were removed by ARACNE. Default: TRUE.
<code>...</code>	Additional arguments passed to 'GENIE3::GENIE3()'

**Value**

A list of data frames representing edge lists. Each list element is an edge list for a specific method.

**Examples**

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_list <- grn_combined(filt.se, regulators=tfs, nTrees=2)
```

**grn\_filter**

*Filter a gene regulatory network based on optimal scale-free topology fit*

**Description**

Filter a gene regulatory network based on optimal scale-free topology fit

**Usage**

```
grn_filter(edgelist, nsplit = 10, bp_param = BiocParallel::SerialParam())
```

**Arguments**

<code>edgelist</code>	A gene regulatory network represented as an edge list.
<code>nsplit</code>	Number of groups in which the edge list will be split. Default: 10.
<code>bp_param</code>	BiocParallel back-end to be used. Default: BiocParallel::SerialParam()

## Details

The edge list will be split in n groups and the scale-free topology fit will be tested for each subset of the edge list. For instance, if an edge list of 10000 rows is used as input, the function will test SFT fit for the top 1000 edges, then top 2000 edges, and so on up to the whole edge list.

## Value

The edge list that best fits the scale-free topology.

## Examples

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_list <- grn_combined(filt.se, regulators=tfs, nTrees=2)
ranked_grn <- grn_average_rank(grn_list)
# split in only 2 groups for demonstration purposes
filtered_edges <- grn_filter(ranked_grn, nsplit=2)
```

**grn\_infer**

*Infer gene regulatory network with one of three algorithms*

## Description

The available algorithms are Context Likelihood of Relatedness (CLR), ARACNE, or GENIE3.

## Usage

```
grn_infer(
  exp,
  regulators = NULL,
  method = c("clr", "aracne", "genie3"),
  estimator_clr = "pearson",
  estimator_aracne = "spearman",
  eps = 0.1,
  remove_zero = TRUE,
  ...
)
```

## Arguments

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a 'SummarizedExperiment' object.
<code>regulators</code>	A character vector of regulators (e.g., transcription factors or miRNAs). All regulators must be included in 'exp'.
<code>method</code>	GRN inference algorithm to be used. One of "clr", "aracne", or "genie3".
<code>estimator_clr</code>	Entropy estimator to be used. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "pearson".

<code>estimator_aracne</code>	
	Entropy estimator to be used. One of "mi.empirical", "mi.mm", "mi.shrink", "mi.sg", "pearson", "spearman", or "kendall". Default: "spearman".
<code>eps</code>	Numeric value indicating the threshold used when removing an edge: for each triplet of nodes (i,j,k), the weakest edge, say (ij), is removed if its weight is below min(ik),(jk) - eps. Default: 0.1.
<code>remove_zero</code>	Logical indicating whether to remove edges whose weight is exactly zero. Default: TRUE
<code>...</code>	Additional arguments passed to 'GENIE3::GENIE3()'.

**Value**

A gene regulatory network represented as an edge list.

**Examples**

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=20, replace=FALSE)
clr <- grn_infer(filt.se, method = "clr", regulators=tfs)
aracne <- grn_infer(filt.se, method = "aracne", regulators=tfs)
# only 2 trees for demonstration purposes
genie3 <- grn_infer(filt.se, method = "genie3", regulators=tfs, nTrees=2)
```

<code>is_singleton</code>	<i>Logical expression to check if gene or gene set is singleton or not</i>
---------------------------	--

**Description**

Logical expression to check if gene or gene set is singleton or not

**Usage**

```
is_singleton(genes, og)
```

**Arguments**

<code>genes</code>	Character containing gene or group of genes to be evaluated.
<code>og</code>	Data frame of 3 columns corresponding to orthogroup, species ID, and gene ID, respectively.

**Value**

Vector of logical values indicating if gene or group of genes is singleton or not.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[is\\_duplicated](#)

**Examples**

```
data(og.zma.osa)
data(filt.se)
genes <- tail(rownames(filt.se), n = 100)
is_singleton(genes, og.zma.osa)
```

---

**modPres\_netrep**

*Calculate module preservation between two expression data sets using NetRep's algorithm*

---

**Description**

Calculate module preservation between two expression data sets using NetRep's algorithm

**Usage**

```
modPres_netrep(
  expList,
  refNet = NULL,
  testNet = NULL,
  nPerm = 1000,
  nThreads = 1
)
```

**Arguments**

<code>expList</code>	List of expression data frames or SummarizedExperiment objects.
<code>refNet</code>	Reference network object returned by the function <code>exp2net</code> .
<code>testNet</code>	Test network object returned by the function <code>exp2net</code> .
<code>nPerm</code>	Number of permutations. Default: 1000
<code>nThreads</code>	Number of threads to be used for parallel computing. Default: 1

**Value**

Output list from `NetRep::modulePreservation` and a message in user's standard output stating which modules are preserved.

**See Also**

[modulePreservation](#)

## Examples

```

set.seed(1)
data(og.zma.osa)
data(zma.se)
data(osa.se)
og <- og.zma.osa
exp_ortho <- exp_genes2orthogroups(expList, og, summarize = "mean")
exp_ortho <- lapply(exp_ortho, function(x) filter_by_variance(x, n=1500))
# Previously calculated SFT powers
powers <- c(13, 15)
gcn.osa <- exp2gcn(exp_ortho$osa, net_type = "signed hybrid",
                     SFTpower = powers[1], cor_method = "pearson")
gcn.zma <- exp2gcn(exp_ortho$zma, net_type = "signed hybrid",
                     SFTpower = powers[2], cor_method = "pearson")
expList <- exp_ortho
ref_net <- gcn.osa
test_net <- gcn.zma
# 10 permutations for demonstration purposes
pres_netrep <- modPres_netrep(expList, ref_net, test_net,
                                nPerm=10, nThreads = 2)

```

**modPres\_WGCNA**

*Calculate module preservation between two expression data sets using WGCNA's algorithm*

## Description

Calculate module preservation between two expression data sets using WGCNA's algorithm

## Usage

```
modPres_WGCNA(expList, ref_net, nPerm = 200)
```

## Arguments

- expList      List of expression data frames or SummarizedExperiment objects.
- ref\_net      Reference network object returned by the function exp2net.
- nPerm        Number of permutations for the module preservation statistics. It must be greater than 1. Default: 200.

## Value

A ggplot object with module preservation statistics.

## Examples

```
set.seed(1)
data(og.zma.osa)
data(zma.se)
data(osa.se)
expList <- list(Zma = zma.se, Osa = osa.se)
og <- og.zma.osa
exp_ortho <- exp_genes2orthogroups(expList, og, summarize = "mean")
exp_ortho <- lapply(exp_ortho, function(x) filter_by_variance(x, n=1500))
# Previously calculated power
powers <- c(13, 15)
gcn.osa <- exp2gcn(exp_ortho$Osa, net_type = "signed hybrid",
                     SFTpower = powers[1], cor_method = "pearson")
expList <- exp_ortho
ref_net <- gcn.osa
# 5 permutations for demonstration purposes
pres_wgcna <- modPres_WGCNA(expList, ref_net, nPerm=5)
```

**module\_enrichment**      *Perform enrichment analysis for coexpression network modules*

## Description

Perform enrichment analysis for coexpression network modules

## Usage

```
module_enrichment(
  net = NULL,
  background_genes,
  annotation,
  column = NULL,
  correction = "BH",
  p = 0.05,
  bp_param = BiocParallel::SerialParam()
)
```

## Arguments

<b>net</b>	List object returned by <code>exp2gcn</code> .
<b>background_genes</b>	Character vector of genes to be used as background for the Fisher's Exact Test.
<b>annotation</b>	Annotation data frame with genes in the first column and functional annotation in the other columns. This data frame can be exported from Biomart or similar databases.

<b>column</b>	Column or columns of annotation to be used for enrichment. Both character or numeric values with column indices can be used. If users want to supply more than one column, input a character or numeric vector. Default: all columns from annotation.
<b>correction</b>	Multiple testing correction method. One of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr" or "none". Default is "BH".
<b>p</b>	P-value threshold. P-values below this threshold will be considered significant. Default is 0.05.
<b>bp_param</b>	BiocParallel back-end to be used. Default: BiocParallel::SerialParam()

**Value**

A data frame containing enriched terms, p-values, gene IDs and module names.

**Author(s)**

Fabricio Almeida-Silva

**Examples**

```
data(filt.se)
data(zma.interpro)
background <- rownames(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
mod_enrich <- module_enrichment(gcn, background, zma.interpro, p=1)
```

**module\_preservation**     *Calculate network preservation between two expression data sets*

**Description**

Calculate network preservation between two expression data sets

**Usage**

```
module_preservation(
  explist,
  ref_net = NULL,
  test_net = NULL,
  algorithm = "netrep",
  nPerm = 1000,
  nThreads = 1
)
```

**Arguments**

expList	List of SummarizedExperiment objects or expression data frames with genes (or orthogroups) in row names and samples in column names.
ref_net	Reference network object returned by the function exp2gcn.
test_net	Test network object returned by the function exp2gcn.
algorithm	Module preservation algorithm to be used. One of 'netrep' (default, permutation-based) or WGCNA.
nPerm	Number of permutations. Default: 1000
nThreads	Number of threads to be used for parallel computing. Default: 1

**Value**

A list containing the preservation statistics (netrep) or a ggplot object with preservation statistics. See WGCNA::modulePreservation or NetRep::modulePreservation for more info.

**Examples**

```
set.seed(1)
data(og.zma.osa)
data(zma.se)
data(osa.se)
og <- og.zma.osa
exp_ortho <- exp_genes2orthogroups(expList, og, summarize = "mean")
exp_ortho <- lapply(exp_ortho, function(x) filter_by_variance(x, n=1500))
# Previously calculated SFT powers
powers <- c(13, 15)
gcn.osa <- exp2gcn(exp_ortho$osa, net_type = "signed hybrid",
                     SFTpower = powers[1], cor_method = "pearson")
gcn.zma <- exp2gcn(exp_ortho$zma, net_type = "signed hybrid",
                     SFTpower = powers[2], cor_method = "pearson")
expList <- exp_ortho
ref_net <- gcn.osa
test_net <- gcn.zma
# 10 permutations for demonstration purposes
pres <- module_preservation(expList, ref_net, test_net, nPerm=10)
```

module_stability	<i>Perform module stability analysis</i>
------------------	--

**Description**

Perform module stability analysis

**Usage**

```
module_stability(exp, net, nRuns = 20)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>net</code>	List object returned by <code>exp2gcn</code> .
<code>nRuns</code>	Number of times to resample. Default is 20.

**Value**

A base plot with the module stability results.

**See Also**

[sampledBlockwiseModules](#),[matchLabels](#),[plotDendroAndColors](#)

**Examples**

```
data(filt.se)
filt <- filt.se[1:100, ] # reducing even further for testing purposes
# The SFT fit was previously calculated and the optimal power was 16
gcn <- exp2gcn(filt, SFTpower = 16, cor_method = "pearson")
# For simplicity, only 2 runs
module_stability(exp = filt, net = gcn, nRuns = 2)
```

*module\_trait\_cor*      *Correlate module eigengenes to trait*

**Description**

Correlate module eigengenes to trait

**Usage**

```
module_trait_cor(
  exp,
  metadata,
  MEs,
  cor_method = "spearman",
  transpose = FALSE,
  palette = "RdYlBu",
  continuous_trait = FALSE,
  cex.lab.x = 0.6,
  cex.lab.y = 0.6,
  cex.text = 0.6
)
```

## Arguments

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>metadata</code>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘exp’ is a ‘SummarizedExperiment’ object, since the function will extract colData.
<code>MEs</code>	Module eigengenes. It is the 2nd element of the result list generated by the function <code>exp2gcn</code> .
<code>cor_method</code>	Method to calculate correlation. One of ‘pearson’, ‘spearman’ or ‘kendall’. Default is ‘spearman’.
<code>transpose</code>	Logical indicating whether to transpose the heatmap of not. Default is FALSE.
<code>palette</code>	RColorBrewer’s color palette to use. Default is “RdYlBu”, a palette ranging from blue to red.
<code>continuous_trait</code>	Logical indicating if trait is a continuous variable. Default is FALSE.
<code>cex.lab.x</code>	Font size for x axis labels. Default: 0.6.
<code>cex.lab.y</code>	Font size for y axis labels. Default: 0.6.
<code>cex.text</code>	Font size for numbers inside matrix. Default: 0.6.

## Details

Significance levels: 1 asterisk: significant at alpha = 0.05. 2 asterisks: significant at alpha = 0.01. 3 asterisks: significant at alpha = 0.001. no asterisk: not significant.

## Value

A data frame with correlation and correlation p-values for each pair of ME and trait along with a heatmap.

## Author(s)

Fabricio Almeida-Silva

## Examples

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
module_trait_cor(filt.se, MEs=gcn$MEs)
```

<code>net_stats</code>	<i>Calculate network statistics</i>
------------------------	-------------------------------------

## Description

Calculate network statistics

## Usage

```
net_stats(
  adj_matrix = NULL,
  net_type = c("gcn", "ppi", "grn"),
  calculate_additional = FALSE
)
```

## Arguments

adj_matrix	Adjacency matrix that represents the network.
net_type	One of "gcn" (gene coexpression network), "ppi" (protein-protein interaction), or "grn" (gene regulatory network).
calculate_additional	Logical indicating whether to calculate additional network statistics (betweenness and closeness). Default is FALSE.

## Value

A list containing the following elements:

- Connectivity
- ScaledConnectivity
- ClusterCoef
- MAR (for gcn only)
- Density
- Centralization
- Heterogeneity (gcn only)
- Diameter
- Betweenness
- Closeness

## See Also

[graph\\_from\\_adjacency\\_matrix](#), [cliques](#), [diameter](#), [estimate\\_betweenness](#), [V](#), [closeness](#), [degree](#), [transitivity](#), [edge\\_density](#), [centr\\_degree](#) [fundamentalNetworkConcepts](#)

## Examples

```
data(filt.se)
set.seed(12)
filt.se <- exp_preprocess(
  filt.se, Zk_filtering = FALSE, variance_filter = TRUE, n = 200
)
gcn <- exp2gcn(
  filt.se, SFTpower = 7, cor_method = "pearson", net_type = "signed hybrid"
)
stats <- net_stats(gcn$adjacency_matrix, net_type = "gcn")
```

---

og.zma.osa

*Orthogroups between maize and rice*

---

## Description

The orthogroups were downloaded from the PLAZA 4.0 Monocots database.

## Usage

```
data(og.zma.osa)
```

## Format

A 3-column data frame with orthogroups, species IDs and gene IDs.

## References

Van Bel, M., Diels, T., Vancaester, E., Kreft, L., Botzki, A., Van de Peer, Y., ... & Vandepoele, K. (2018). PLAZA 4.0: an integrative resource for functional, evolutionary and comparative plant genomics. Nucleic acids research, 46(D1), D1190-D1196.

## Examples

```
data(og.zma.osa)
```

---

`osa.se`*Rice gene expression data from Shin et al., 2021.*

---

**Description**

Filtered expression data in transcripts per million (TPM) from Shin et al., 2021. Genes with TPM values <5 in more than 60 were removed to reduce package size. The expression data and associated sample metadata are stored in a SummarizedExperiment object.

**Usage**

```
data(osa.se)
```

**Format**

An object of class `SummarizedExperiment`

**References**

Shin, J., Marx, H., Richards, A., Vaneechoutte, D., Jayaraman, D., Maeda, J., ... & Roy, S. (2021). A network-based comparative framework to study conservation and divergence of proteomes in plant phylogenies. *Nucleic Acids Research*, 49(1), e3-e3.

**Examples**

```
data(osa.se)
```

---

`parse_orthofinder`*Parse orthogroups identified by OrthoFinder*

---

**Description**

This function converts the orthogroups file named **Orthogroups.tsv** to a 3-column data frame that can be interpreted by BioNERO.

**Usage**

```
parse_orthofinder(file_path = NULL)
```

**Arguments**

<code>file_path</code>	Path to Orthogroups/Orthogroups.tsv file generated by OrthoFinder.
------------------------	--

**Value**

A 3-column data frame with orthogroups, species IDs and gene IDs, respectively.

**Author(s)**

Fabricio Almeida-Silva

**Examples**

```
path <- system.file("extdata", "Orthogroups.tsv", package = "BioNERO")
og <- parse_orthofinder(path)
```

---

PC_correction	<i>Apply Principal Component (PC)-based correction for confounding artifacts</i>
---------------	--

---

**Description**

Apply Principal Component (PC)-based correction for confounding artifacts

**Usage**

```
PC_correction(exp, verbose = FALSE)
```

**Arguments**

- |         |  |
|---------|--|
| exp     | A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment‘ object. |
| verbose | Logical indicating whether to display progress messages or not. Default: FALSE.                                      |

**Value**

Corrected expression data frame or ‘SummarizedExperiment‘ object.

**Author(s)**

Fabricio Almeida-Silva

**References**

- Parsana, P., Ruberman, C., Jaffe, A. E., Schatz, M. C., Battle, A., & Leek, J. T. (2019). Addressing confounding artifacts in reconstruction of gene co-expression networks. *Genome biology*, 20(1), 1-6.

**See Also**

[num.sv,sva\\_network](#)

**Examples**

```
data(zma.se)
exp <- filter_by_variance(zma.se, n=500)
exp <- PC_correction(exp)
```

**plot\_dendro\_and\_colors**

*Plot dendrogram of genes and modules*

### Description

Plot dendrogram of genes and modules

### Usage

`plot_dendro_and_colors(gcn)`

### Arguments

`gcn` List object returned by `exp2gcn`.

### Value

A base plot with the gene dendrogram and modules.

### Examples

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
plot_dendro_and_colors(gcn)
```

**plot\_dendro\_and\_cons\_colors**

*Plot dendrogram of genes and consensus modules*

### Description

Plot dendrogram of genes and consensus modules

### Usage

`plot_dendro_and_cons_colors(consensus)`

### Arguments

`consensus` Consensus network returned by `consensus_modules`.

### Value

A base plot with the gene dendrogram and modules.

## Examples

```
set.seed(12)
data(zma.se)
filt.zma <- filter_by_variance(zma.se, n=500)
zma.set1 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
zma.set2 <- filt.zma[, sample(colnames(filt.zma), size=20, replace=FALSE)]
list.sets <- list(zma.set1, zma.set2)
# SFT power previously identified with consensus_SFT_fit()
cons_mod <- consensus_modules(list.sets, power = c(11, 13),
                                cor_method = "pearson")
plot_dendro_and_cons_colors(cons_mod)
```

---

## plot\_eigengene\_network

*Plot eigengene network*

---

## Description

Plot eigengene network

## Usage

```
plot_eigengene_network(gcn)
```

## Arguments

gcn                  List object returned by exp2gcn.

## Value

A base plot with the eigengene network

## Examples

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
plot_eigengene_network(gcn)
```

**plot\_expression\_profile***Plot expression profile of given genes across samples***Description**

Plot expression profile of given genes across samples

**Usage**

```
plot_expression_profile(
  genes,
  exp,
  metadata,
  plot_module = TRUE,
  net,
  modulename
)
```

**Arguments**

<code>genes</code>	Character vector containing a subset of genes from which edges will be extracted. It can be ignored if <code>plot_module</code> is TRUE.
<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>metadata</code>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘exp’ is a ‘SummarizedExperiment’ object, since the function will extract colData.
<code>plot_module</code>	Logical indicating whether to plot a whole module or not. If set to FALSE, genes must be specified.
<code>net</code>	List object returned by <code>exp2net</code> .
<code>modulename</code>	Name of the module to plot.

**Value**

A ggplot object showing the expression profile of some genes across all samples.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[melt](#)

## Examples

```
data(zma.se)
data(filt.se)
genes <- rownames(filt.se)
plot_expression_profile(genes=genes, exp=zma.se, plot_module=FALSE)
```

**plot\_gcn**

*Plot gene coexpression network from edge list*

## Description

Plot gene coexpression network from edge list

## Usage

```
plot_gcn(
  edgelist_gcn,
  net,
  color_by = "module",
  hubs = NULL,
  show_labels = "tophubs",
  top_n_hubs = 5,
  interactive = FALSE,
  dim_interactive = c(600, 600)
)
```

## Arguments

<code>edgelist_gcn</code>	Data frame containing the edge list for the GCN. The edge list can be generated with <code>get_edge_list()</code> .
<code>net</code>	List object returned by <code>exp2net</code> .
<code>color_by</code>	How should nodes be colored? It must be either "module" (nodes will have the colors of their modules) or a 2-column data frame containing genes in the first column and a custom gene annotation in the second column. Default: "module".
<code>hubs</code>	Data frame containing hub genes in the first column, their modules in the second column, and intramodular connectivity in the third column.
<code>show_labels</code>	Character indicating which nodes will be labeled. One of "all", "allhubs", "tophubs", or "none". Default: tophubs.
<code>top_n_hubs</code>	Number of top hubs to be labeled. It is only valid if <code>show_labels</code> equals "tophubs". Default is 5.
<code>interactive</code>	Logical indicating whether the network should be interactive or not. Default is FALSE.
<code>dim_interactive</code>	Numeric vector with width and height of window for interactive plotting. Default: c(600,600).

**Value**

A ggplot object.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

```
simplify,as\_data\_frame,gorder igraph\_to\_networkD3,forceNetwork geom\_edges,geom\_nodes,geom\_nodetext,them
```

**Examples**

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
gcn_edges <- get_edge_list(gcn, module="brown", filter=TRUE,
                             method="min_cor")
hubs <- get_hubs_gcn(filt.se, gcn)
p <- plot_gcn(gcn_edges, gcn, hubs = hubs)
```

**plot\_grn**

*Plot gene regulatory network from edge list*

**Description**

Plot gene regulatory network from edge list

**Usage**

```
plot_grn(
  edgelist_grn,
  show_labels = "tophubs",
  top_n_hubs = 5,
  interactive = FALSE,
  layout = igraph::with_kk,
  arrow.gap = 0.01,
  ranked = TRUE,
  dim_interactive = c(600, 600)
)
```

**Arguments**

- |                           |   |
|---------------------------|---|
| <code>edgelist_grn</code> | Data frame containing the edge list for the GRN network. First column is the TF and second column is the target gene. All other columns are interpreted as edge attributes. |
| <code>show_labels</code>  | Character indicating which nodes will be labeled. One of "all", "allhubs", "tophubs", or "none".  |

top_n_hubs	Number of top hubs to be labeled. It is only valid if show_labels equals "top_hubs". Default is 5.
interactive	Logical indicating whether the network should be interactive or not. Default is FALSE.
layout	igraph function for the network layout. One of with_dh, with_drl, with_gem, with_lgl, with_fr, with_graphopt, with_kk and with_mds. Default is with_kk.
arrow.gap	Numeric indicating the distance between nodes and arrows. Default is 0.2.
ranked	Logical indicating whether to treat third column of the edge list (edge weights) as ranked values. Default: TRUE.
dim_interactive	Numeric vector with width and height of window for interactive plotting. Default: c(600,600).

**Value**

A ggplot object containing the network.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[igraph\\_to\\_networkD3](#), [forceNetwork](#) [geom\\_edges](#), [geom\\_nodes](#), [geom\\_nodetext](#), [theme\\_blank](#), [geom\\_nodetext\\_repel](#) [new\\_scale](#)

**Examples**

```
data(filt.se)
tfs <- sample(rownames(filt.se), size=50, replace=FALSE)
grn_edges <- grn_infer(filt.se, method ="clr", regulators = tfs)
p <- plot_grn(grn_edges, ranked=FALSE)
```

**plot\_heatmap**

*Plot heatmap of hierarchically clustered sample correlations or gene expression*

**Description**

Plot heatmap of hierarchically clustered sample correlations or gene expression

**Usage**

```
plot_heatmap(
  exp,
  col_metadata = NA,
  row_metadata = NA,
  cor_method = "spearman",
  type = "samplecor",
  palette = NULL,
  log_trans = FALSE,
  cluster_rows = TRUE,
  cluster_cols = TRUE,
  show_rownames = FALSE,
  show_colnames = TRUE,
  scale = "none",
  fontsize = 9,
  cutree_rows = NA,
  cutree_cols = NA,
  ...
)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>col_metadata</code>	A data frame containing sample names in row names and sample annotation in the subsequent columns. The maximum number of columns is 2 to ensure legends can be visualized. Ignored if ‘exp’ is a ‘SummarizedExperiment’ object, since the function will extract colData. Default: NA.
<code>row_metadata</code>	A data frame containing gene IDs in row names and gene functional classification in the first column. Only one column is allowed to ensure legends can be visualized. Default: NA.
<code>cor_method</code>	Correlation method. One of ‘spearman’ or ‘pearson’. Default is ‘spearman’.
<code>type</code>	Type of heatmap to plot. One of ‘samplecor’ (sample correlations) or ‘expr’. Default: ‘samplecor’.
<code>palette</code>	RColorBrewer palette to use. Default is “Blues” for sample correlation heatmap and “YlOrRd” for gene expression heatmap.
<code>log_trans</code>	Logical indicating whether to log transform the expression data or not. Default: FALSE.
<code>cluster_rows</code>	Logical indicating whether to cluster rows or not. Default: TRUE.
<code>cluster_cols</code>	Logical indicating whether to cluster columns or not. Default: TRUE.
<code>show_rownames</code>	Logical indicating whether to show row names or not. Default: FALSE.
<code>show_colnames</code>	Logical indicating whether to show column names or not. Default is TRUE.
<code>scale</code>	Character indicating if values should be centered and scaled in rows, columns, or none. One of ‘row’, ‘column’, or ‘none’. Default: ‘none’.
<code>fontsize</code>	Base font size for the plot.

```
cutree_rows      Number of clusters into which rows are divided. Default: NA (no division).
cutree_cols      Number of clusters into which columns are divided. Default: NA (no division).
...              Additional arguments to be passed to ComplexHeatmap::pheatmap().
```

### Value

A heatmap of sample correlations or gene expression.

### Author(s)

Fabricio Almeida-Silva

### See Also

[RColorBrewer](#)

### Examples

```
data(filt.se)
plot_heatmap(filt.se)
```

---

## plot\_ngenes\_per\_module

*Plot number of genes per module*

---

### Description

Plot number of genes per module

### Usage

```
plot_ngenes_per_module(net = NULL)
```

### Arguments

net List object returned by exp2gcn.

### Value

A ggplot object with a bar plot of gene number in each module.

### Examples

```
data(filt.se)
gcn <- exp2gcn(filt.se, SFTpower = 18, cor_method = "pearson")
plot_ngenes_per_module(gcn)
```

**plot\_PCA***Plot Principal Component Analysis (PCA) of samples***Description**

Plot Principal Component Analysis (PCA) of samples

**Usage**

```
plot_PCA(exp, metadata, log_trans = FALSE, PCs = "1x2", size = 2)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
<code>metadata</code>	A data frame containing sample names in row names and sample annotation in the first column. Ignored if ‘exp‘ is a ‘SummarizedExperiment‘ object, since the function will extract colData.
<code>log_trans</code>	Logical. If TRUE, the expression data frame will be log transformed with $\log_2(\text{exp}+1)$ .
<code>PCs</code>	Principal components to be plotted on the x-axis and y-axis, respectively. One of "1x2", "1x3" or "2x3". Default is "1x2".
<code>size</code>	Numeric indicating the point size. Default is 2.

**Value**

A ggplot object with the PCA plot.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[ggplot](#)

**Examples**

```
data(zma.se)
plot_PCA(zma.se, log_trans = TRUE)
```

---

plot_ppi	<i>Plot protein-protein interaction network from edge list</i>
----------	--

---

## Description

Plot protein-protein interaction network from edge list

## Usage

```
plot_ppi(  
  edgelist_int,  
  color_by = "community",  
  clustering_method = igraph::cluster_infomap,  
  show_labels = "tophubs",  
  top_n_hubs = 5,  
  interactive = FALSE,  
  add_color_legend = TRUE,  
  dim_interactive = c(600, 600)  
)
```

## Arguments

edgelist_int	Data frame containing the edge list for the PPI network. First column is the protein 1 and second column is the protein 2. All other columns are interpreted as edge attributes.
color_by	How should nodes be colored? It must be either "community" or a 2-column data frame containing proteins in the first column and a custom annotation in the second column. If "community", a clustering algorithm will be applied. Default: "community".
clustering_method	igraph function to be used for community detection. Available functions are cluster_infomap, cluster_edge_betweenness, cluster_fast_greedy, cluster_walktrap, cluster_springlass, cluster_leading_eigen, cluster_louvain, and cluster_label_prop. Default is cluster_infomap.
show_labels	Character indicating which nodes will be labeled. One of "all", "allhubs", "tophubs", or "none".
top_n_hubs	Number of top hubs to be labeled. It is only valid if show_labels equals "tophubs". Default is 5.
interactive	Logical indicating whether the network should be interactive or not. Default is FALSE.
add_color_legend	Logical indicating whether to add a color legend for nodes. Default: TRUE.
dim_interactive	Numeric vector with width and height of window for interactive plotting. Default: c(600,600).

**Value**

A ggplot object.

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[as\\_data\\_frame](#), [degree](#), [simplify](#), [gorder](#), [igraph\\_to\\_networkD3](#), [forceNetwork](#)

**Examples**

```
ppi_edges <- igraph:::get.edgelist(igraph::barabasi.game(n=50, directed=FALSE))
p <- plot_ppi(ppi_edges, add_color_legend = FALSE)
```

---

**q\_normalize**

*Quantile normalize the expression data*

---

**Description**

Quantile normalize the expression data

**Usage**

```
q_normalize(exp)
```

**Arguments**

<code>exp</code>	A gene expression data frame with genes in row names and samples in column names.
------------------	---

**Value**

Expression matrix with normalized values

**Examples**

```
data(zma.se)
exp <- SummarizedExperiment::assay(zma.se)
norm_exp <- q_normalize(exp)
```

---

remove_nonexp	<i>Remove genes that are not expressed based on a user-defined threshold</i>
---------------	--

---

## Description

Remove genes that are not expressed based on a user-defined threshold

## Usage

```
remove_nonexp(  
  exp,  
  method = "median",  
  min_exp = 1,  
  min_percentage_samples = 0.25  
)
```

## Arguments

- exp** A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment’ object.
- method** Criterion to filter non-expressed genes out. One of "mean", "median", "percentage", or "allsamples". Default is "median".
- min\_exp** If method is 'mean', 'median', or 'allsamples', the minimum value for a gene to be considered expressed. If method is 'percentage', the minimum value each gene must have in at least n percent of samples to be considered expressed.
- min\_percentage\_samples** In case the user chooses 'percentage' as method, expressed genes must have expression  $\geq$  min\_exp in at least this percentage. Values must range from 0 to 1.

## Value

Filtered gene expression data frame or ‘SummarizedExperiment’ object.

## Author(s)

Fabricio Almeida-Silva

## See Also

[rowMedians](#) [goodSamplesGenes](#)

## Examples

```
data(zma.se)  
filt_exp <- remove_nonexp(zma.se, min_exp = 5)
```

---

`replace_na`*Remove missing values in a gene expression data frame*

---

**Description**

Remove missing values in a gene expression data frame

**Usage**

```
replace_na(exp, replaceby = 0)
```

**Arguments**

- |                        |  |
|------------------------|--|
| <code>exp</code>       | A gene expression data frame with genes in row names and samples in column names or a ‘SummarizedExperiment‘ object. |
| <code>replaceby</code> | What to use instead of NAs. One of 0 or ‘mean’. Default is 0.  |

**Value**

Gene expression data frame or ‘SummarizedExperiment‘ object with all NAs replaced according to the argument ‘replaceby’.

**Author(s)**

Fabricio Almeida-Silva

**Examples**

```
data(zma.se)
exp <- replace_na(zma.se)
sum(is.na(exp))
```

---

`SFT_fit`*Pick power to fit network to a scale-free topology*

---

**Description**

Pick power to fit network to a scale-free topology

**Usage**

```
SFT_fit(exp, net_type = "signed", rsquared = 0.8, cor_method = "spearman")
```

**Arguments**

exp	A gene expression data frame with genes in row names and samples in column names or a 'SummarizedExperiment' object.
net_type	Network type. One of 'signed', 'signed hybrid' or 'unsigned'. Default is signed.
rsquared	R squared cutoff. Default is 0.8.
cor_method	Correlation method. One of "pearson", "biweight" or "spearman". Default is "spearman".

**Value**

A list containing:

- powerOptimal power based on scale-free topology fit
- plotA ggplot object displaying main statistics of the SFT fit test

**Author(s)**

Fabricio Almeida-Silva

**See Also**

[pickSoftThreshold](#)

**Examples**

```
data(filt.se)
sft <- SFT_fit(filt.se, cor_method = "pearson")
```

ZKfiltering

*Filter outlying samples based on the standardized connectivity (Zk) method*

**Description**

Filter outlying samples based on the standardized connectivity (Zk) method

**Usage**

```
ZKfiltering(exp, zk = -2, cor_method = "spearman")
```

**Arguments**

exp	A gene expression data frame with genes in row names and samples in column names or a 'SummarizedExperiment' object.
zk	Standardized connectivity threshold. Default is -2.
cor_method	Correlation method. One of "pearson", "biweight" or "spearman". Default is "spearman".

**Value**

Filtered gene expression data frame or ‘SummarizedExperiment’ object.

**Author(s)**

Fabricio Almeida-Silva

**References**

Oldham, M. C., Langfelder, P., & Horvath, S. (2012). Network methods for describing sample relationships in genomic datasets: application to Huntington’s disease. *BMC systems biology*, 6(1), 1-18.

**See Also**

[adjacency](#)

**Examples**

```
data(zma.se)
filt_exp <- ZKfiltering(zma.se)
```

---

zma.interpro

*Maize Interpro annotation*

---

**Description**

Interpro protein domain annotation retrieved from the PLAZA Monocots 4.0 database. Only genes included in zma.se are present in this subset.

**Usage**

```
data(zma.interpro)
```

**Format**

A 2-column data frame containing gene IDs and their associated Interpro annotations.

**References**

Van Bel, M., Diels, T., Vancaester, E., Kreft, L., Botzki, A., Van de Peer, Y., ... & Vandepoele, K. (2018). PLAZA 4.0: an integrative resource for functional, evolutionary and comparative plant genomics. *Nucleic acids research*, 46(D1), D1190-D1196.

**Examples**

```
data(zma.interpro)
```

---

zma.se

*Maize gene expression data from Shin et al., 2021.*

---

## Description

Filtered expression data in transcripts per million (TPM) from Shin et al., 2021. Genes with TPM values <5 in more than 60 were removed to reduce package size. The expression data and associated sample metadata are stored in a SummarizedExperiment object.

## Usage

```
data(zma.se)
```

## Format

An object of class SummarizedExperiment

## References

Shin, J., Marx, H., Richards, A., Vaneechoutte, D., Jayaraman, D., Maeda, J., ... & Roy, S. (2021). A network-based comparative framework to study conservation and divergence of proteomes in plant phylogenies. *Nucleic Acids Research*, 49(1), e3-e3.

## Examples

```
data(zma.se)
```

---

zma.tfs

*Maize transcription factors*

---

## Description

Transcription factors and their families were downloaded from PlantTFDB 4.0.

## Usage

```
data(zma.tfs)
```

## Format

A data frame with gene IDs of TFs and their associated families.

## References

Jin, J., Tian, F., Yang, D. C., Meng, Y. Q., Kong, L., Luo, J., & Gao, G. (2016). PlantTFDB 4.0: toward a central hub for transcription factors and regulatory interactions in plants. *Nucleic acids research*, gkw982.

**Examples**

```
data(zma.tfs)
```

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