

# Package ‘SEMgraph’

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**Title** Network Analysis and Causal Inference Through Structural Equation Modeling

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**Description** Estimate networks and causal relationships in complex systems through Structural Equation Modeling. This package also includes functions to import, weight, manipulate, and fit biological network models within the Structural Equation Modeling framework described in Grassi M, Palluzzi F, Tarantino B (2022) <[doi:10.1093/bioinformatics/btac567](https://doi.org/10.1093/bioinformatics/btac567)>.

**URL** <https://github.com/fernandoPalluzzi/SEMgraph>

**Depends** igraph (>= 1.3.0), lavaan (>= 0.5-23), R (>= 4.0)

**Imports** boot (>= 1.3-25), corpcor (>= 1.6.9), dagitty (>= 0.3-0), flip (>= 2.5.0), gdata (>= 2.18.0), ggm (>= 2.3), glmnet (>= 2.0-18), graph (>= 1.56.0), leaps (>= 3.1), mgcv (>= 1.8-39), pbapply (>= 1.4-3), pcalg (>= 2.7-6), protoclust (>= 1.6.3), RBGL (>= 1.68.0), Rgraphviz (>= 2.22.0)

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**Maintainer** Barbara Tarantino <[barbara.tarantino01@universitadipavia.it](mailto:barbara.tarantino01@universitadipavia.it)>

**Repository** CRAN

**Author** Mario Grassi [aut],  
Fernando Palluzzi [aut],  
Barbara Tarantino [aut, cre]

**R topics documented:**

activeModule . . . . .	3
alsData . . . . .	5
ancestry . . . . .	6
clusterGraph . . . . .	7
clusterScore . . . . .	8
colorGraph . . . . .	10
corr2graph . . . . .	12
cplot . . . . .	14
dagitty2graph . . . . .	15
extractClusters . . . . .	16
factor.analysis . . . . .	19
gplot . . . . .	20
graph2dag . . . . .	22
graph2dagitty . . . . .	23
graph2lavaan . . . . .	24
gsa.extract . . . . .	24
kegg . . . . .	26
kegg.pathways . . . . .	27
lavaan2graph . . . . .	28
localCI.test . . . . .	29
mergeNodes . . . . .	30
model.learn . . . . .	32
modelSearch . . . . .	35
orientEdges . . . . .	38
pairwiseMatrix . . . . .	39
parameterEstimates . . . . .	40
pathFinder . . . . .	40
properties . . . . .	42
resizeGraph . . . . .	43
sachs . . . . .	45
sem.mediation . . . . .	46
SEMace . . . . .	48
SEMbap . . . . .	50
SEMdag . . . . .	53
SEMdci . . . . .	55
SEMgsa . . . . .	57
SEMpath . . . . .	59
SEMrun . . . . .	60
SEMtree . . . . .	64
Shipley.test . . . . .	67
summary.GGM . . . . .	69
summary.RICF . . . . .	69
weightGraph . . . . .	70

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activeModule	<i>Active module identification</i>
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## Description

Uses different information flow and tree-based strategies for identifying active modules (e.g., disease modules), including a perturbed subset of nodes and edges. Function scalability enables graph reduction at both pathway and entire interactome scales.

## Usage

```
activeModule(
  graph,
  type,
  seed,
  eweight = "none",
  alpha = 0.05,
  top = 100,
  limit = 10000,
  ...
)
```

## Arguments

graph	An igraph object.
type	Module identification method. If type = "kou", the Steiner tree algorithm will be applied. If type = "usp", the resulting graph will be the union of all significant shortest paths. If type = "rwr", the random walk with restart algorithm will be enabled. Finally, if type = "hdi", the heat diffusion algorithm is used.
seed	Either a user-defined vector containing seed node names or one among: "pvlm", "proto", or "qi", corresponding to the seed name attribute yielded by <a href="#">weightGraph</a> .
eweight	Edge weight type derived from <a href="#">weightGraph</a> or from user-defined distances. This option determines the weight-to-distance transform. If set to "none" (default), edge weights will be set to 1. If eweight = "kegg", repressing interactions (-1) will be set to 1 (maximum distance), neutral interactions (0) will be set to 0.5, and activating interactions (+1) will be set to 0 (minimum distance). If eweight = "zsign", all significant interactions will be set to 0 (minimum distance), while non-significant ones will be set to 1. If eweight = "pvalue", weights (p-values) will be transformed to the inverse of negative base-10 logarithm. If eweight = "custom", the algorithm will use the distance measure specified by the user as "weight" edge attribute.
alpha	Significance level to assess shortest paths significance, when type is "usp". By default, alpha = 0.05.
top	Number of top nodes for the "rwr" and "hdi" algorithms. The output subgraph is induced by the top-n ranking nodes. By default, top = 100 (i.e., the top-100 of nodes are selected).

limit	An integer value corresponding to the number of graph edges. If type = "usp", beyond this limit, multicore computation is enabled to reduce the computational burden. By default, limit = 10000.
...	Currently ignored.

## Details

Graph filtering algorithms include:

1. "kou", the Steiner tree connecting a set of seed nodes, using the algorithm suggested by Kou et al. (1981);
2. "usp", generates a subnetwork as the union of the significant (P-value < alpha) shortest paths between the seeds set;
3. "rwr", Random Walk with Restart, a wrapper for random.walk function of the R package diffusr;
4. "hdi", Heat Diffusion algorithm, a wrapper for heat.diffusion function of the R package diffusr.

## Value

An active module, an igraph object with colored nodes (seed = "green", and connector = "white").

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

Palluzzi F, Grassi M (2021). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. <arXiv:2103.08332>

Kou L, Markowsky G, Berman L (1981). A fast algorithm for Steiner trees. Acta Informatica, 15(2): 141-145. <<https://doi.org/10.1007/BF00288961>>

Simon Dirmeier (2018). diffusr: Network Diffusion Algorithms. R package version 0.1.4. <<https://CRAN.R-project.org/package=diffusr/>>

## Examples

```
# Graph weighting
G <- weightGraph(graph = sachs$graph, data = sachs$pkc, group = sachs$group,
  method = "r2z",
  seed = c(0.05, 0.5, 0.5))

# RWR algorithm, seeds and edge P-values as weights
R1 <- activeModule(graph = G, type = "kou", seed = "pvlm", eweight = "pvalue")
R2 <- activeModule(graph = G, type = "kou", seed = "proto", eweight = "pvalue")
R3 <- activeModule(graph = G, type = "kou", seed = "qi", eweight = "pvalue")

# Graphs
old.par <- par(no.readonly = TRUE)
```

```

par(mfrow=c(2,2), mar=rep(2, 4))
plot(G, layout = layout.circle, main = "input graph")
box(col = "gray")
plot(R1, layout = layout.circle, main = "lm P-value (alpha = 0.05)")
box(col = "gray")
plot(R2, layout = layout.circle, main = "prototype (h = 0.5)")
box(col = "gray")
plot(R3, layout = layout.circle, main = "closeness (q = 0.5)")
box(col = "gray")
par(old.par)

```

alsData

*Amyotrophic Lateral Sclerosis (ALS) dataset*

## Description

Expression profiling through high-throughput sequencing (RNA-seq) of 139 ALS patients and 21 healthy controls (HCs), from Tam et al. (2019).

## Usage

```
alsData
```

## Format

alsData is a list of 4 objects:

1. "graph", ALS graph as the largest connected component of the "Amyotrophic lateral sclerosis (ALS)" pathway from KEGG database;
2. "exprs", a matrix of 160 rows (subjects) and 318 columns (genes) extracted from the original 17695. This subset includes genes from KEGG ALS signaling pathway, MAPK signaling pathway, and Protein processing in endoplasmic reticulum pathway, needed to run SEMgraph examples. Raw data from the GEO dataset GSE124439 (Tam et al., 2019) were pre-processed applying batch effect correction, using the sva R package (Leek et al., 2012), to remove data production center and brain area biases. Using multidimensional scaling-based clustering, ALS-specific and an HC-specific clusters were generated. Misclassified samples were black-listed and removed from the current dataset;
3. "group", a binary group vector of 139 ALS subjects (1) and 21 healthy controls (0);
4. "details", a data.frame reporting information about included and blacklisted samples.

## Source

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE124439/>

## References

Tam OH, Rozhkov NV, Shaw R, Kim D et al. (2019). Postmortem Cortex Samples Identify Distinct Molecular Subtypes of ALS: Retrotransposon Activation, Oxidative Stress, and Activated Glia. *Cell Rep* 29(5):1164-1177.e5. <<https://doi.org/10.1016/j.celrep.2019.09.066>>

Jeffrey T. Leek, W. Evan Johnson, Hilary S. Parker, Andrew E. Jaffe, and John D. Storey (2012). The sva package for removing batch effects and other unwanted variation in high-throughput experiments. *Bioinformatics*. Mar 15; 28(6): 882-883. <<https://doi.org/10.1093/bioinformatics/bts034>>

## Examples

```
alsData$graph
dim(alsData$exprs)
table(alsData$group)
```

---

ancestry

*Node ancestry utilities*

---

## Description

Get ancestry for a collection of nodes in a graph. These functions are wrappers for the original SEMID R package.

## Usage

```
ancestors(g, nodes)

descendants(g, nodes)

parents(g, nodes)

siblings(g, nodes)
```

## Arguments

<code>g</code>	An igraph object.
<code>nodes</code>	the nodes in the graph of which to get the ancestry.

## Value

a sorted vector of nodes.

## References

Rina Foygel Barber, Mathias Drton and Luca Weihs (2019). SEMID: Identifiability of Linear Structural Equation Models. R package version 0.3.2. <<https://CRAN.R-project.org/package=SEMID/>>

**Examples**

```
# Get all ancestors
an <- V(sachs$graph)[ancestors(sachs$graph, "Erk")]; an

# Get parents
pa <- V(sachs$graph)[parents(sachs$graph, "PKC")]; pa

# Get descendants
de <- V(sachs$graph)[descendants(sachs$graph, "PKA")]; de

# Get siblings
sib <- V(sachs$graph)[siblings(sachs$graph, "PIP3")]; sib
```

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clusterGraph	<i>Topological graph clustering</i>
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---

**Description**

Topological graph clustering methods.

**Usage**

```
clusterGraph(graph, type = "wtc", HM = "none", size = 5, verbose = FALSE, ...)
```

**Arguments**

graph	An igraph object.
type	Topological clustering methods. If type = "tahc", network modules are generated using the tree agglomerative hierarchical clustering method (Yu et al., 2015). Other non-tree clustering methods from <a href="#">igraph</a> package include: "wtc" (default value; walktrap community structure with short random walks), "ebc" (edge betweenness clustering), "fgc" (fast greedy method), "lbc" (label propagation method), "lec" (leading eigenvector method), "loc" (multi-level optimization), "opc" (optimal community structure), "sgc" (springlass statistical mechanics).
HM	Hidden model type. Enables the visualization of the hidden model, gHM. If set to "none" (default), no gHM igraph object is saved. For each defined hidden module: (i) if HM = "LV", a latent variable (LV) will be defined as common unknown cause acting on cluster nodes; (ii) if HM = "CV", cluster nodes will be considered as regressors of a latent composite variable (CV); (iii) if HM = "UV", an unmeasured variable (UV) is defined, where source nodes of the module (i.e., in-degree = 0) act as common regressors influencing the other nodes via an unmeasured variable (see also <a href="#">clusterScore</a> ).
size	Minimum number of nodes per module. By default, a minimum number of 5 nodes is required.

verbose            A logical value. If FALSE (default), the gHM igraph will not be plotted to screen, saving execution time (they will be returned in output anyway).

...                Currently ignored.

**Value**

If HM is not "none" a list of 2 objects is returned:

1. "gHM", subgraph containing hidden modules as an igraph object;
2. "membership", cluster membership vector for each node.

If HM is "none", only the cluster membership vector is returned.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

Fortunato S, Hric D. Community detection in networks: A user guide (2016). Phys Rep; 659: 1-44. <<https://dx.doi.org/10.1016/j.physrep.2016.09.002>>

Yu M, Hillebrand A, Tewarie P, Meier J, van Dijk B, Van Mieghem P, Stam CJ (2015). Hierarchical clustering in minimum spanning trees. Chaos 25(2): 023107. <<https://doi.org/10.1063/1.4908014>>

**See Also**

[clusterScore](#), [cplot](#)

**Examples**

```
# Clustering ALS graph with WTC method and LV model
G <- properties(alsData$graph)[[1]]
clv <- clusterGraph(graph = G, type = "wtc", HM = "LV")
gplot(clv$gHM, l = "fdp")
table(clv$membership)
```

---

clusterScore	<i>Module scoring</i>
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---

**Description**

Generate factor scores, principal component scores, or projection scores of latent, composite, and unmeasured variable modules, respectively, and fit them with an exogenous group effect.



**Usage**

```
clusterScore(
  graph,
  data,
  group,
  HM = "LV",
  type = "wtc",
  size = 5,
  verbose = FALSE,
  ...
)
```

**Arguments**

graph	An igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes.
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects.
HM	Hidden model type. For each defined hidden module: (i) if HM = "LV", a latent variable (LV) will be defined as common unknown cause acting on cluster nodes; (ii) if HM = "CV", cluster nodes will be considered as regressors of a latent composite variable (CV); (iii) if HM = "UV", an unmeasured variable (UV) model will be generated for each module, where source nodes (i.e., in-degree = 0) act as common regressors influencing the other nodes via an unmeasured variable. By default, HM is set to "LV" (i.e., the latent variable model).
type	Graph clustering method. If type = "tahc", network modules are generated using the tree agglomerative hierarchical clustering method (Yu et al., 2015). Other non-tree clustering methods from igraph package include: "wtc" (default value; walktrap community structure with short random walks), "ebc" (edge betweenness clustering), "fgc" (fast greedy method), "lbc" (label propagation method), "lec" (leading eigenvector method), "loc" (multi-level optimization), "opc" (optimal community structure), "sgc" (springlass statistical mechanics). By default, the "wtc" method is used.
size	Minimum number of nodes per hidden module. By default, a minimum number of 5 nodes is required.
verbose	A logical value. If TRUE, intermediate graphs will be displayed during the execution. In addition, a reduced graph with clusters as nodes will be fitted and showed to screen (see also <a href="#">mergeNodes</a> ). By default, verbose = FALSE.
...	Currently ignored.

**Value**

A list of 3 objects:

1. "fit", hidden module fitting as a lavaan object;

2. "membership", hidden module nodes membership; `clusterGraph` function;
3. "dataHM", data matrix with cluster scores in first columns.

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

### References

Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38 (20), 4829–4830 <<https://doi.org/10.1093/bioinformatics/btac567>>

### See Also

See `clusterGraph` and `cplot` for graph clustering.

### Examples

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

C <- clusterScore(graph = alsData$graph, data = als.npn,
                  group = alsData$group,
                  HM = "LV",
                  type = "wtc",
                  verbose = FALSE)

summary(C$fit)
head(C$dataHM)
table(C$membership)
```

---

colorGraph

*Vertex and edge graph coloring on the base of fitting*

---

### Description

Add vertex and edge color attributes to an igraph object, based on a fitting results data.frame generated by `SEMrun`.

### Usage

```
colorGraph(
  est,
  graph,
  group,
  method = "none",
  alpha = 0.05,
```

```

vcolor = c("lightblue", "white", "pink"),
ecolor = c("royalblue3", "gray50", "red2"),
ewidth = c(1, 2),
...
)

```

## Arguments

<code>est</code>	A data.frame of estimated parameters and p-values, derived from the <code>fit</code> object returned by <a href="#">SEMrun</a> . As an alternative, the user may provide a "gest" or "dest" data.frame generated by <a href="#">SEMrun</a> .
<code>graph</code>	An igraph object.
<code>group</code>	group A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects.
<code>method</code>	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method is set to "none" (i.e., no multiple test correction).
<code>alpha</code>	Significance level for node and edge coloring (by default, $\alpha = 0.05$ ).
<code>vcolor</code>	A vector of three color names. The first color is given to nodes with P-value < alpha and beta < 0, the third color is given to nodes with P-value < alpha and beta > 0, and the second is given to nodes with P-value > alpha. By default, <code>vcolor = c("lightblue", "white", "pink")</code> .
<code>ecolor</code>	A vector of three color names. The first color is given to edges with P-value < alpha and regression coefficient < 0, the third color is given to edges with P-value < alpha and regression coefficient > 0, and the second is given to edges with P-value > alpha. By default, <code>vcolor = c("blue", "gray50", "red2")</code> .
<code>ewidth</code>	A vector of two values. The first value refers to the basic edge width (i.e., edges with P-value > alpha), while the second is given to edges with P-value < alpha. By default <code>ewidth = c(1, 2)</code> .
<code>...</code>	Currently ignored.

## Value

An igraph object with vertex and edge color and width attributes.

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## Examples

```

# Model fitting: node perturbation
sem1 <- SEMrun(graph = alsData$graph, data = alsData$exprs,
               group = alsData$group,
               fit = 1)
est1 <- parameterEstimates(sem1$fit)

```

```

# Model fitting: edge perturbation
sem2 <- SEMrun(graph = alsData$graph, data = alsData$exprs,
               group = alsData$group,
               fit = 2)
est20 <- subset(parameterEstimates(sem2$fit), group = 1)[, -c(4, 5)]
est21 <- subset(parameterEstimates(sem2$fit), group = 2)[, -c(4, 5)]

# Graphs
g <- alsData$graph
x <- alsData$group

old.par <- par(no.readonly = TRUE)
par(mfrow=c(2,2), mar=rep(1,4))
gplot(colorGraph(est = est1, g, group = x, method = "BH"),
      main = "vertex differences")
gplot(colorGraph(est = sem2$dest, g, group = NULL),
      main = "edge differences")
gplot(colorGraph(est = est20, g, group = NULL),
      main = "edges for group = 0")
gplot(colorGraph(est = est21, g, group = NULL),
      main = "edges for group = 1")
par(old.par)

```

---

corr2graph

*Correlation matrix to graph*


---

## Description

Convert a correlation matrix to an igraph object.

## Usage

```
corr2graph(R, n, type = "marg", method = "none", alpha = 0.05, ...)
```

## Arguments

R	Correlation matrix.
n	Sample size (i.e., the number of subjects).
type	Graph building method. If type is either "marg" or "cond", marginal or conditional correlation tests will be used, respectively. If type = "mst", input correlations are converted to distances and a minimum spanning tree is generated from the distance matrix, using Prim's algorithm (Prim, 1957). If type = "tmfg", a triangulate maximally graph is generated from the given correlation matrix (Massara et al., 2016).

method	Multiple testing correction method. One of the values available in <code>p.adjust</code> . By default, method = "none" (i.e., no multiple test correction). See <code>p.adjust</code> for other correction methods.
alpha	Significance level used to compute the correlation threshold. By default, alpha = 0.05.
...	Currently ignored.

**Value**

An igraph object.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

- Palluzzi F, Grassi M (2021). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. <arXiv:2103.08332>
- Massara GP, Di Matteo T and Aste T (2009). Network Filtering for Big Data: Triangulated Maximally Filtered Graph. Journal of complex Networks, 5(2): 161–178. <<https://doi.org/10.1093/comnet/cnw015>>
- Prim RC (1957). Shortest connection networks and some generalizations. Bell System Technical Journal, 36(6):1389–1401. <<https://doi.org/10.1002/j.1538-7305.1957.tb01515.x>>

**Examples**

```
# Graphs creation
C1 <- corr2graph(R = cor(log(sachs$pkc)), n = nrow(sachs$pkc),
                type = "marg",
                method = "BH")
C2 <- corr2graph(R = cor(log(sachs$pkc)), n = nrow(sachs$pkc),
                type = "cond",
                method = "BH")
C3 <- corr2graph(R = cor(log(sachs$pkc)), n = nrow(sachs$pkc),
                type = "mst",
                method = "BH")
C4 <- corr2graph(R = cor(log(sachs$pkc)), n = nrow(sachs$pkc),
                type = "tmfg",
                method = "BH")

# Graphs plots
old.par <- par(no.readonly = TRUE)
par(mfrow=c(2,2), mar= rep(2, 4))
plot(C1, layout=layout.circle, main= "marg"); box(col="gray")
plot(C2, layout=layout.circle, main= "cond"); box(col="gray")
plot(C3, layout=layout.circle, main= "mst"); box(col="gray")
plot(C4, layout=layout.circle, main= "tmfg"); box(col="gray")
par(old.par)
```

cplot

*Subgraph mapping***Description**

Map groups of nodes onto an input graph, based on a membership vector.

**Usage**

```
cplot(graph, membership, l = layout.auto, map = FALSE, verbose = FALSE, ...)
```

**Arguments**

graph	An igraph object.
membership	Cluster membership vector for each node.
l	graph layout. One of the <a href="#">igraph</a> layouts. If this argument is ignored, an automatic layout will be applied.
map	A logical value. Visualize cluster mapping over the input graph. If FALSE (default), visualization will be disabled. For large graphs, visualization may take long.
verbose	A logical value. If FALSE (default), the processed graphs will not be plotted to screen, saving execution time (they will be returned in output anyway).
...	Currently ignored.

**Value**

The list of clusters and cluster mapping as igraph objects.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**See Also**

[clusterGraph](#), [clusterScore](#)

**Examples**

```
# Clustering ALS graph with WTC method
G <- alsData$graph
membership <- clusterGraph(graph = G, type = "wtc")
cplot(G, membership, map = TRUE, verbose = FALSE)
cplot(G, membership, map = FALSE, verbose = TRUE)
# The list of cluster graphs !
cg <- cplot(G, membership); cg
```

---

dagitty2graph	<i>Graph conversion from dagitty to igraph</i>
---------------	------------------------------------------------

---

**Description**

Convert a dagitty object to a igraph object.

**Usage**

```
dagitty2graph(dagi, verbose = FALSE, ...)
```

**Arguments**

dagi	A graph as a dagitty object ("dag" or "pdag").
verbose	A logical value. If TRUE, the output graph is shown (for graph2dagitty only). This argument is FALSE by default.
...	Currently ignored.

**Value**

An igraph object.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**Examples**

```
# Conversion from igraph to dagitty (and viceversa)
dagi <- graph2dagitty(sachs$graph, verbose = TRUE)
graph <- dagitty2graph(dagi, verbose = TRUE)
```

---

extractClusters	<i>Cluster extraction utility</i>
-----------------	-----------------------------------

---

## Description

Extract and fit clusters from an input graph.

Extract and fit clusters from an input graph.

## Usage

```
extractClusters(
  graph,
  data,
  membership,
  group = NULL,
  n = 2000,
  fitting.method = "lavaan",
  size = 5,
  ...
)
```

```
extractClusters(
  graph,
  data,
  membership,
  group = NULL,
  n = 2000,
  fitting.method = "lavaan",
  size = 5,
  ...
)
```

## Arguments

graph	Input network as an igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
membership	A vector of cluster membership IDs. The membership vector can be generated using the <a href="#">clusterScore</a> , <a href="#">clusterGraph</a> , or <a href="#">model.learn</a> functions.
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. Group specification enables node perturbation testing. By default, group = NULL.
n	Number of randomization replicates (default = 2000), for permutation flip or bootstrap sampling, if algo = "ricf".



fitting.method	MLE method used for SEM fitting. If algo = "lavaan" (default), the SEM will be fitted using the NLMINB solver from lavaan R package, with standard errors derived from the expected Fisher information matrix. If algo = "ricf", the model is fitted via residual iterative conditional fitting (RICF; Drton et al. 2009). If algo = "cggm", model fitting is based on constrained Gaussian Graphical Modeling (CGGM; Hastie et al. 2009, p. 446).
size	Minimum size for a cluster to be fitted (default = 5). Use size = 0 to enforce model fitting for every cluster.
...	Currently ignored.
map	Logical value. If TRUE, the plot of the input graph (coloured by cluster membership) will be generated along with independent module plots. If the input graph is very large, plotting could be computationally intensive (by default, map = FALSE).
verbose	Logical value. If TRUE, a plot will be showed for each cluster.

### Value

(i) clusters: list clusters as igraph objects; (ii) fit: list of fitting results for each cluster as a lavaan object; (iii) dfc: data.frame of summary results

A list of three objects:

1. "clusters", A list of clusters as igraph objects;
2. "estimates", A data.frame containing model estimates for each cluster;
3. "fit.indices", A data.frame containing fitting indices for each cluster.

### Author(s)

Fernando Palluzzi <fernando.palluzzi@gmail.com>

### References

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**See Also**

See [clusterScore](#) and [clusterGraph](#) for network clustering. See [model.learn](#) for data-driven model learning and clustering. See [SEMrun](#) for general model fitting.

**Examples**

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

adjdata <- SEMbap(alsData$graph, als.npn)$data

# Clusters creation
clusters <- extractClusters(graph = alsData$graph, data = adjdata)
print(clusters$dfc)
head(parameterEstimates(clusters$fit$HM1))
head(parameterEstimates(clusters$fit$HM2))
head(parameterEstimates(clusters$fit$HM4))
gplot(clusters$clusters$HM2)

# Map cluster on the input graph
g <- alsData$graph
c <- clusters$clusters$HM2
V(g)$color <- ifelse(V(g)$name %in% V(c)$name, "gold", "white")
gplot(g)
```

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

# Cluster creation
clusters <- clusterScore(graph = alsData$graph, data = als.npn,
                        group = alsData$group,
                        type = "wtc")

# Clusters extraction and fitting with the RICF algorithm
fit <- extractClusters(graph = alsData$graph, data = als.npn,
                      membership = clusters$membership,
                      group = alsData$group,
                      fitting.method = "ricf")

# Extracting differentially regulated nodes
DRNs <- rownames(fit$estimates)[fit$estimates$pvalue < 0.05]
length(DRNs)

# Map cluster 1 on the input graph
g <- alsData$graph
```

```
c <- fit$clusters[[1]]
V(g)$color <- ifelse(V(g)$name %in% V(c)$name, "gold", "white")
gplot(g)
```

---

factor.analysis

*Factor analysis for high dimensional data*


---

## Description

Wrapper for Factor Analysis with potentially high dimensional variables implement in the "cate" R package (Author: Jingshu Wang [aut], Qingyuan Zhao [aut, cre] Maintainer: Qingyuan Zhao <qz280@cam.ac.uk>) that is optimized for the high dimensional problem where the number of samples  $n$  is less than the number of variables  $p$ .

## Usage

```
factor.analysis(Y, r = 1, method = "pc")
```

## Arguments

<code>Y</code>	data matrix, a $n \times p$ matrix
<code>r</code>	number of factors (default, $r=1$ )
<code>method</code>	algorithm to be used, "pc" (default) or "ml"

## Details

The two methods extracted from "cate" are quasi-maximum likelihood (ml), and principal component analysis (pc). The ml is iteratively solved the EM algorithm using the PCA solution as the initial value. See Bai and Li (2012) for more details.

## Value

a list of objects

**Gamma** estimated factor loadings

**Z** estimated latent factors

**Sigma** estimated noise variance matrix

## References

Jushan Bai and Kunpeng Li (2012). Statistical Analysis of Factor Models of High Dimension. The Annals of Statistics, 40 (1), 436-465 <<https://doi.org/10.1214/11-AOS966>>

Jingshu Wang and Qingyuan Zhao (2020). cate: High Dimensional Factor Analysis and Confounder Adjusted Testing and Estimation. R package version 1.1.1. <<https://CRAN.R-project.org/package=cate>>

**Examples**

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

## pc
pc<- factor.analysis(Y = als.npn, r = 2, method = "pc")
head(pc$Gamma)
head(pc$Z)
head(pc$Sigma)

## ml
ml <- factor.analysis(Y = als.npn, r = 2, method = "ml")
head(ml$Gamma)
head(ml$Z)
head(ml$Sigma)
```

---

gplot

*Graph plotting with renderGraph*

---

**Description**

Wrapper for function renderGraph of the R package Rgraphviz.

**Usage**

```
gplot(
  graph,
  l = "dot",
  main = "",
  cex.main = 1,
  font.main = 1,
  color.txt = "black",
  fontsize = 16,
  cex = 0.6,
  shape = "circle",
  color = "gray70",
  lty = 1,
  lwd = 1,
  w = "auto",
  h = "auto",
  psize = 80,
  ...
)
```

**Arguments**

<code>graph</code>	An igraph or graphNEL object.
<code>l</code>	Any layout supported by Rgraphviz. It can be one among: "dot" (default), "neato", "circo", "fdp", "osage", "twopi".
<code>main</code>	Plot main title (by default, no title is added).
<code>cex.main</code>	Main title size (default = 1).
<code>font.main</code>	Main title font (default = 1). Available options are: 1 for plain text, 2 for bold, 3 for italics, 4 for bold italics, and 5 for symbol.
<code>color.txt</code>	Node text color (default = "black").
<code>fontsize</code>	Node text size (default = 16).
<code>cex</code>	Another argument to control node text size (default = 0.6).
<code>shape</code>	Node shape (default = "circle").
<code>color</code>	Node border color (default = "gray70").
<code>lty</code>	Node border outline (default = 1). Available options include: 0 for blank, 1 for solid line, 2 for dashed, 3 for dotted, 4 for dotdash, 5 for longdash, and 6 for twodash.
<code>lwd</code>	Node border thickness (default = 1).
<code>w</code>	Manual node width (default = "auto").
<code>h</code>	Manual node height (default = "auto").
<code>psize</code>	Automatic node size (default = 80).
<code>...</code>	Currently ignored.

**Value**

`gplot` returns invisibly the graph object produced by Rgraphviz

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**Examples**

```
gplot(sachs$graph, main = "input graph")

sem <- SEMrun(sachs$graph, sachs$pkc)
gplot(sem$graph, main = "output graph")
```

graph2dag

*Convert directed graphs to directed acyclic graphs (DAGs)***Description**

Remove cycles and bidirected edges from a directed graph.

**Usage**

```
graph2dag(graph, data, bap = FALSE, time.limit = Inf, ...)
```

**Arguments**

graph	A directed graph as an igraph object.
data	A data matrix with subjects as rows and variables as columns.
bap	If TRUE, a bow-free acyclic path (BAP) is returned (default = FALSE).
time.limit	CPU time for the computation, in seconds (default = Inf).
...	Currently ignored.

**Details**

The conversion is performed firstly by removing bidirected edges and then the data matrix is used to compute edge P-values, through marginal correlation testing (see [weightGraph](#), r-to-z method). When a cycle is detected, the edge with highest P-value is removed, breaking the cycle. If the bap argument is TRUE, a BAP is generated merging the output DAG and the bidirected edges from the input graph.

**Value**

A DAG as an igraph object.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**Examples**

```
dag <- graph2dag(graph = sachs$graph, data = log(sachs$pkc))
old.par <- par(no.readonly = TRUE)
par(mfrow=c(1,2), mar=rep(1, 4))
gplot(sachs$graph, main = "Input graph")
gplot(dag, main = "Output DAG")
par(old.par)
```

---

graph2dagitty	<i>Graph conversion from igraph to dagitty</i>
---------------	------------------------------------------------

---

## Description

Convert an igraph object to a dagitty object.

## Usage

```
graph2dagitty(graph, graphType = "dag", verbose = FALSE, ...)
```

## Arguments

graph	A graph as an igraph or as an adjacency matrix.
graphType	character, is one of "dag" (default) or "pdag". DAG can contain the directed (->) and bi-directed (<->) edges, while PDAG can contain the edges: ->, <->, and the undirected edges (-) that represent edges whose direction is not known.
verbose	A logical value. If TRUE, the output graph is shown (for graph2dagitty only). This argument is FALSE by default.
...	Currently ignored.

## Value

A dagitty object.

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## Examples

```
# Graph as an igraph object to dagitty object
G <- graph2dagitty(sachs$graph)
plot(dagitty::graphLayout(G))
```

---

graph2lavaan	<i>Graph to lavaan model</i>
--------------	------------------------------

---

**Description**

Convert an igraph object to a model (lavaan syntax).

**Usage**

```
graph2lavaan(graph, nodes = V(graph)$name, ...)
```

**Arguments**

graph	A graph as an igraph object.
nodes	Subset of nodes to be included in the model. By default, all the input graph nodes will be included in the output model.
...	Currently ignored.

**Value**

A model in lavaan syntax.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**Examples**

```
# Graph (igraph object) to structural model in lavaan syntax
model <- graph2lavaan(sachs$graph)
cat(model, "\n")
```

---

gsa.extract	<i>Gene set analysis extraction utility</i>
-------------	---------------------------------------------

---

**Description**

Extract differentially regulated nodes (DRNs) and interaction network(s) from the [SEMgsa](#) output. The interaction network is obtained by the union of all the perturbed pathways.



**Usage**

```
gsa.extract(
  gsa,
  reference = reactome.pathways,
  lcc = TRUE,
  min.size = 5,
  max.size = 300,
  min.drns = 2,
  act.alpha = 0.05,
  inh.alpha = 0.05,
  adj.alpha = 0.05,
  ...
)
```

**Arguments**

<code>gsa</code>	Gene Set Analysis object generated by <a href="#">SEMgsa</a> .
<code>reference</code>	Reference list of biological pathways. Every element of the list must be an <code>igraph</code> object. <code>SEMgraph</code> provides both Reacrome and KEGG pathways list. By default, <code>reactome.pathways</code> is used.
<code>lcc</code>	A logical value. If <code>TRUE</code> (default) only the largest connected component of the resulting network is considered.
<code>min.size</code>	Minimum required pathway size (default = 5).
<code>max.size</code>	Maximum allowed pathway size (default = 300).
<code>act.alpha</code>	Pathway activation significance level (default = 0.05).
<code>inh.alpha</code>	Pathway inhibition significance level (default = 0.05).
<code>adj.alpha</code>	Global cutoff over the adjusted perturbation p-value (default = 0.05).
<code>...</code>	Currently ignored.
<code>min.degs</code>	Minimum required number of DRNs (default = 2).

**Value**

A list of two objects:

1. "model", The output model as an `igraph` object;
2. "clusters", A list containing model clustering results. It includes tree more objects:
  - "membership", A vector reporting cluster membership for each node;
  - "fit", hidden model fitting as a lavaan object;
  - "dataHM", data matrix containing cluster scores (hidden variables) alongside the original data variables.

**Author(s)**

Fernando Palluzzi <fernando.palluzzi@gmail.com>

## References

Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38 (20), 4829–4830 <<https://doi.org/10.1093/bioinformatics/btac567>>

## See Also

[SEMgsa](#)

## Examples

```
# Nonparanormal(npn) transformation

library(huge)
als.npn <- huge.npn(alsData$exprs)

# Selection of FTD-ALS pathways from kegg.pathways.Rdata

paths.name <- c("MAPK signaling pathway",
               "Protein processing in endoplasmic reticulum",
               "Endocytosis",
               "Wnt signaling pathway",
               "Neurotrophin signaling pathway",
               "Amyotrophic lateral sclerosis")

j <- which(names(kegg.pathways) %in% paths.name)

GSA <- SEMgsa(kegg.pathways[j], als.npn, alsData$group,
             method = "bonferroni", alpha = 0.05,
             n_rep = 1000)

gsa.pert <- gsa.extract(GSA, kegg.pathways)
```

---

kegg

*KEGG interactome*

---

## Description

Interactome generated by merging KEGG pathways extracted using the R0ntoTools R package (update: November, 2021).

## Usage

kegg

**Format**

"kegg" is an igraph network object of 5934 nodes and 77158 edges (41122 directed and 3164/2 = 1582 bidirected) corresponding to the union of 225 KEGG pathways.

**Source**

<https://www.genome.jp/kegg/>

**References**

Kanehisa M, Goto S (1999). KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acid Research 28(1): 27-30. <<https://doi.org/10.1093/nar/27.1.29>>

Calin Voichita, Sahar Ansari and Sorin Draghici (2021). ROntoTools: R Onto-Tools suite. R package version 2.20.0.

**Examples**

```
# KEGG graph
summary(kegg)

# KEGG degrees of freedom
vcount(kegg)*(vcount(kegg) - 1)/2 - ecount(kegg)
```

---

kegg.pathways

*KEGG pathways*

---

**Description**

KEGG pathways extracted using the ROntoTools R package (update: November, 2021).

**Usage**

```
kegg.pathways
```

**Format**

"kegg.pathways" is a list of 225 igraph objects corresponding to the KEGG pathways.

**Source**

<https://www.genome.jp/kegg/>

**References**

Kanehisa M, Goto S (1999). KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acid Research 28(1): 27-30. <<https://doi.org/10.1093/nar/27.1.29>>

Calin Voichita, Sahar Ansari and Sorin Draghici (2021). ROntoTools: R Onto-Tools suite. R package version 2.20.0.

## Examples

```
library(igraph)

# KEGG pathways
names(kegg.pathways)

i<-which(names(kegg.pathways)=="Type II diabetes mellitus");i
ig<- kegg.pathways[[i]]
summary(ig)
V(ig)$name
E(ig)$weight

gplot(ig, l="fdp", psize=50, main=names(kegg.pathways[i]))
```

---

lavaan2graph

*lavaan model to graph*


---

## Description

Convert a model, specified using lavaan syntax, to a graph object in either igraph or dagitty format.

## Usage

```
lavaan2graph(model, directed = TRUE, psi = TRUE, verbose = FALSE, ...)
```

## Arguments

<code>model</code>	Model specified using lavaan syntax.
<code>directed</code>	Logical value. If TRUE (default), edge directions from the model will be preserved. If FALSE, the resulting graph will be undirected.
<code>psi</code>	Logical value. If TRUE (default) covariances will be converted into bidirected graph edges. If FALSE, covariances will be excluded from the output graph.
<code>verbose</code>	Logical value. If TRUE, a plot of the output graph will be generated. For large graphs, this could significantly increase computation time. If FALSE (default), graph plotting will be disabled.
<code>...</code>	Currently ignored.

## Value

An igraph object.

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## Examples

```
# Writing path diagram in lavaan syntax

model<-'
#path model
Jnk ~ PKA + PKC
P38 ~ PKA + PKC
Akt ~ PKA + PIP3
Erk ~ PKA + Mek
Mek ~ PKA + PKC + Raf
Raf ~ PKA + PKC
PKC ~ PIP2 + Plcg
PIP2 ~ PIP3 + Plcg
Plcg ~ PIP3
#PKA ~ 1
#PIP3 ~ 1

# (co)variances
# PIP2 ~~ PIP3
'

# Graph with covariances
G0 <- lavaan2graph(model, psi = TRUE)
plot(G0, layout = layout.circle)

# Graph without covariances
G1 <- lavaan2graph(model, psi = FALSE)
plot(G1, layout = layout.circle)
```

---

localCI.test

---

*Conditional Independence (CI) local tests of an acyclic graph*


---

## Description

P-values of one minimal testable implication (with the smallest possible conditioning set) is returned per missing edge given an acyclic graph (DAG or BAP) using the function [impliedConditionalIndependencies](#) plus the function [localTests](#) from package `dagitty`. Without assuming any particular dependence structure, the p-values of every CI test, in a DAG (BAP), is then combined using the Bonferroni's statistic in an overall test of the fitted model,  $B = K \cdot \min(p_1, \dots, p_K)$ , as reviewed in Vovk & Wang (2020).

## Usage

```
localCI.test(graph, data, bap = FALSE, limit = 10000, verbose = TRUE, ...)
```

**Arguments**

graph	A directed graph as an igraph object.
data	A data matrix with subjects as rows and variables as columns.
bap	If TRUE, the input graph is transformed in a BAP, if FALSE (default) the input graph is reduced in a DAG.
limit	An integer value corresponding to the number of missing edges of the extracted acyclic graph. Beyond this limit, switch to Shipley's C-test (Shipley 2000) is enabled to reduce the computational burden. By default, limit = 10000.
verbose	If TRUE, LocalCI results will be showed to screen (default = TRUE).
...	Currently ignored.

**Value**

A list of three objects: (i) "dag": the DAG used to perform the localCI test (ii) "mseps": the list of all m-separation tests over missing edges in the input graph and (iii) "mtest": the overall Bonferroni's P-value.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

Vovk V, Wang R (2020). Combining p-values via averaging. *Biometrika* 107(4): 791-808. <<https://doi.org/10.1093/biomet/as>>  
 Shipley B (2000). A new inferential test for path models based on DAGs. *Structural Equation Modeling*, 7(2): 206-218. <[https://doi.org/10.1207/S15328007SEM0702\\_4](https://doi.org/10.1207/S15328007SEM0702_4)>

**Examples**

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

sem <- SEMrun(alsData$graph, als.npn)
B_test <- localCI.test(sem$graph, als.npn, verbose = TRUE)
```

---

mergeNodes

*Graph nodes merging by a membership attribute*


---

**Description**

Merge groups of graph nodes using hierarchical clustering with prototypes derived from [protoclust](#) or custom membership attribute (e.g., cluster membership derived from [clusterGraph](#)).

**Usage**

```
mergeNodes(
  graph,
  data,
  h = 0.5,
  membership = NULL,
  HM = NULL,
  verbose = FALSE,
  ...
)
```

**Arguments**

graph	network as an igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes. If membership is not NULL, is currently ignored, data = NULL.
h	Cutting the minimax clustering at height, $h = 1 - \text{abs}(\text{cor}(j,k))$ , yielding a merged node (and a reduced data set) in which every node in the cluster has correlation of at least $\text{cor}(j,k)$ with the prototype node. By default, $h = 0.5$ , i.e. $\text{cor}(j,k) = 0.5$ .
membership	Cluster membership. A vector of cluster membership identifiers as numeric values, where vector names correspond to graph node names. By default, membership = NULL.
HM	Hidden cluster label. If membership is derived from clusterGraph: HM = "LV", a latent variable (LV) will be defined as common unknown cause acting on cluster nodes. If HM = "CV", cluster nodes will be considered as regressors of a latent composite variable (CV). Finally, if HM = "UV", an unmeasured variable (UV) is defined, where source nodes of the module (i.e., in-degree = 0) act as common regressors influencing the other nodes via an unmeasured variable. By default, HM = NULL
verbose	A logical value. If FALSE (default), the merged graphs will not be plotted to screen.
...	Currently ignored.

**Details**

Hierarchical clustering with prototypes (or Minmax linkage) is unique in naturally associating a node (the prototypes) with every interior node of the dendrogram. Thus, for each merge we have a single representative data point for the resulting cluster (Bien, Tibshirani, 2011). These prototypes can be used to greatly enhance the interpretability of merging nodes and data reduction for SEM fitting.

**Value**

A list of 2 objects is returned:

1. "gLM", A graph with merged nodes as an igraph object;
2. "membership", cluster membership vector for each node.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

Bien J, Tibshirani R (2011). Hierarchical Clustering With Prototypes via Minimax Linkage. Journal of the American Statistical Association 106(495): 1075-1084. <doi:10.1198/jasa.2011.tm10183>

**See Also**

[clusterGraph](#)

**Examples**

```
# Gene memberships with prototypes with h=0.5
G <- properties(alsData$graph)[[1]]
M <- mergeNodes(G, data = alsData$exprs, h = 0.5, verbose=TRUE)

# Gene memberships with EBC method and size=10
m <- clusterGraph(G, type = "ebc", size = 10)
M <- mergeNodes(G, membership = m, HM = "LV", verbose=TRUE)

# Gene memberships defined by user
c1 <- c("5894", "5576", "5567", "572", "598")
c2 <- c("6788", "84152", "2915", "836", "5530")
c3 <- c("5603", "6300", "1432", "5600")
m <- c(rep(1,5), rep(2,5), rep(3,4))
names(m) <- c(c1, c2, c3)
M <- mergeNodes(G, membership = m, HM = "CV", verbose=TRUE)
```

---

model.learn

*Data-driven model learning*

---

**Description**

Data-driven model structure learning based on the linear order provided by the input graph.

**Usage**

```
model.learn(
  graph,
  data,
  group = NULL,
  method = "TO",
  beta = 0,
  dag.only = FALSE,
  cluster = "wtc",
```



```

    hidden = "LV",
    min.clust = 5,
    edge.weights = TRUE
  )

```

### Arguments

graph	Input network as an igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. If NULL (default), group influence will not be considered.
method	Linear order method. If method = "TO", the topological order of the input DAG is enabled (default). If codemethod = "TD" a data-driven top-down minimum conditional variance method is performed.
beta	Numeric value corresponding to the minimum LASSO beta coefficient needed for a new interaction to be retained in the final model (default = 0). Increasing beta values decrease output model density and generally increase the number of output clusters.
dag.only	A logical value. If FALSE (default), inferred connections will be added to the input graph. If TRUE, only inferred connections will be retained in the output. In this case, the output network will be a directed acyclic graph. See <a href="#">SEMDag</a> for details.
cluster	Graph clustering method. Basic clustering methods are taken from the igraph package, including: "wtc" (default value; walktrap community structure with short random walks), "ebc" (edge betweenness clustering), "fgc" (fast greedy method), "lbc" (label propagation method), "lec" (leading eigenvector method), "loc" (multi-level optimization), "opc" (optimal community structure), "sgc" (springlass statistical mechanics). If type = "talc", network modules are generated using the tree agglomerative hierarchical clustering method (Yu et al., 2015). Clustering can be disabled by setting cluster = NULL.
hidden	Hidden model type. For each defined hidden module: (i) if HM = "LV", a latent variable (LV) will be defined as common unknown cause acting on cluster nodes; (ii) if HM = "CV", cluster nodes will be considered as regressors of a latent composite variable (CV); (iii) if HM = "UV", an unmeasured variable (UV) model will be generated for each module, where source nodes (i.e., in-degree = 0) act as common regressors influencing the other nodes via an unmeasured variable. By default, HM is set to "LV" (i.e., the latent variable model).
min.clust	Minimum size for a cluster to be fitted (default = 5). Ignored if cluster = NULL.
edge.weights	Logical value. If TRUE, the resulting network is weighted using the r2z method (see <a href="#">weightGraph</a> ).
...	Currently ignored.

**Value**

A list of two objects:

1. "model", The output model as an igraph object;
2. "clusters", A list containing model clustering results. It includes three more objects:
  - "membership", A vector reporting cluster membership for each node;
  - "fit", hidden model fitting as a lavaan object;
  - "dataHM", data matrix containing cluster scores (hidden variables) alongside the original data variables.

**Author(s)**

Fernando Palluzzi <fernando.palluzzi@gmail.com>

**References**

- Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38 (20), 4829–4830 <<https://doi.org/10.1093/bioinformatics/btac567>>
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**See Also**

[SEMdag](#) for data-driven DAG learning details.

**Examples**

```
library(huge)
als.npn <- huge.npn(alsData$exprs)

# Model learning
model <- model.learn(graph = alsData$graph, data = als.npn,
                     group = alsData$group, method = "TO",
                     beta = 0)
```

---

modelSearch

*Optimal model search strategies*


---

## Description

Four model search strategies are implemented combining SEMdag(), SEMbap(), and resizeGraph() functions. All strategies estimate a new graph by 1) adjusting (BAP deconfounding) the the data matrix and 2) re-sizing the output DAG.

## Usage

```
modelSearch(
  graph,
  data,
  gnet = NULL,
  d = 2,
  search = "basic",
  beta = 0,
  method = "BH",
  alpha = 0.05,
  limit = 30000,
  verbose = FALSE,
  ...
)
```

## Arguments

graph	Input graph as an igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
gnet	Reference directed network used to validate and import nodes and interactions.
d	Maximum allowed geodesic distance for directed or undirected shortest path search. A distance $d = 0$ disables shortest path search (fixed in <code>search = "basic"</code> ), while $d = 1$ (fixed in <code>search = "direct"</code> ) only search for directed links (i.e., no mediators are allowed). A distance $d > 1$ (defaults to $d = 2$ for "outer" and "inner" strategies), will search for shortest paths with at most $d - 1$ mediators between nodes sharing a significant estimated interaction. Connectors are imported from the reference interactome, as specified by the argument <code>gnet</code> . If the edges of the reference interactome are weighted by P-value, as defined by the <code>E(gnet)\$pv</code> attribute, the shortest path with the smallest sum of weights will be chosen (e.g., see <a href="#">weightGraph</a> for graph weighting options).
search	Search strategy. Four model search strategies are available: <ul style="list-style-type: none"> <li>"outer". The estimated DAG is re-sized using <a href="#">resizeGraph</a> to find new indirect paths (i.e., inferred directed connections that may hide new mediators). New interactions and connectors will be searched and imported from</li> </ul>

the reference network (argument `gnet`, see above). Both DAG and extended graph complexity can be controlled with  $\beta > 0$  and  $d > 1$  arguments, respectively. The term "outer" means that new model mediator variables are imported from an external resource (i.e., the reference network).

- "inner". This strategy is analogous to the "outer" one, but disables external mediator search. In other words, new indirect paths are generated by adding new interactions of the input model, so that mediators will be nodes already present in the input graph. The reference network is still used to validate new model paths. Also in this case,  $\beta > 0$  and  $d > 1$  are used.
- "direct". The input graph structure is improved through direct (i.e., adjacent) link search, followed by interaction validation and import from the reference network, with no mediators (i.e.,  $d = 1$ ).
- "basic" (default). While the previous strategies rely on the input graph and the reference network to integrate knowledge to the final model, the "basic" strategy is data-driven. The input graph is needed to define the topological order. The argument `gnet` is set to `NULL` (i.e., no reference network is needed) and argument  $d = 0$ . Model complexity can be still controlled by setting  $\beta > 0$ .

<code>beta</code>	Numeric value. Minimum absolute LASSO $\beta$ coefficient for a new interaction to be retained in the estimated DAG backbone. Lower $\beta$ values correspond to more complex DAGs. By default, $\beta$ is set to 0 (i.e., maximum complexity).
<code>method</code>	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, <code>method</code> is set to "BH" (i.e., Benjamini-Hochberg multiple test correction).
<code>alpha</code>	Significance level for false discovery rate (FDR) used for local d-separation tests. This argument is used to control data de-correlation. A higher $\alpha$ level includes more hidden covariances, thus considering more sources of confounding. If $\alpha = 0$ , data de-correlation is disabled. By default, $\alpha = 0.05$ .
<code>limit</code>	An integer value corresponding to the number of missing edges of the extracted acyclic graph. Beyond this limit, multicore computation is enabled to reduce the computational burden.
<code>verbose</code>	If <code>TRUE</code> , it shows intermediate graphs during the execution (not recommended for large graphs).
<code>...</code>	Currently ignored.

## Details

Search strategies can be ordered by decreasing conservativeness respect to the input graph, as: "direct", "inner", "outer", and "basic". The first three strategies are knowledge-based, since they require an input graph and a reference network, together with data, for knowledge-assisted model improvement. The last one does not require any reference and the output model structure will be data-driven. Output model complexity can be limited using arguments  $d$  and  $\beta$ . While  $d$  is fixed to 0 or 1 in "basic" or "direct", respectively; we suggest starting with  $d = 2$  (only one mediator) for the other two strategies. For knowledge-based strategies, we suggest to start with  $\beta = 0$ . Then,  $\beta$  can be relaxed (0 to  $< 0.1$ ) to improve model fitting, if needed. Since data-driven models can be complex, we suggest to start from  $\beta = 0$  when using the "basic" strategy. The  $\beta$  value

can be relaxed until a good model fit is obtained. Argument `alpha` determines the extent of data adjustment: lower `alpha` values for FDR correction correspond to a smaller number of significant confounding factors, hence a weaker correction (default `alpha = 0.05`).

### Value

The output model as well as the adjusted dataset are returned as a list of 2 objects:

- "graph", the output model as an `igraph` object;
- "data", the adjusted dataset.

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

### Examples

```
# Comparison among different model estimation strategies

library(huge)
als.npn <- huge.npn(alsData$exprs)

# Models estimation
m1 <- modelSearch(graph = alsData$graph, data = als.npn, gnet = kegg,
  search = "direct", beta = 0, alpha = 0.05)
m2 <- modelSearch(graph = alsData$graph, data = als.npn, gnet = kegg,
  d = 2, search = "inner", beta = 0, alpha = 0.05)
m3 <- modelSearch(graph = alsData$graph, data = als.npn, gnet = kegg,
  d = 2, search = "outer", beta = 0, alpha = 0.05)
m4 <- modelSearch(graph = alsData$graph, data = als.npn, gnet = NULL,
  search = "basic", beta = 0.1, alpha = 0.05)

# Graphs
old.par <- par(no.readonly = TRUE)
par(mfrow=c(2,2), mar= rep(1,4))
gplot(m1$graph, main = "direct graph")
gplot(m2$graph, main = "inner graph")
gplot(m3$graph, main = "outer graph")
gplot(m4$graph, main = "basic graph")
par(old.par)
```

---

orientEdges	<i>Assign edge orientation of an undirected graph</i>
-------------	-------------------------------------------------------

---

## Description

Assign edge orientation of an undirected graph through a given reference directed graph. The vertex (color) and edge (color, width and weight) attributes of the input undirected graph are preserved in the output directed graph.

## Usage

```
orientEdges(ug, dg, ...)
```

## Arguments

ug	An undirected graph as an igraph object.
dg	A directed reference graph.
...	Currently ignored.

## Value

A directed graph as an igraph object.

## Examples

```
# Graphs definition
G0 <- as.undirected(sachs$graph)

# Reference graph-based orientation
G1 <- orientEdges(ug = G0, dg = sachs$graph)

# Graphs plotting
old.par <- par(no.readonly = TRUE)
par(mfrow=c(1,2), mar=rep(2,4))
plot(G0, layout=layout.circle, main = "Input undirected graph")
plot(G1, layout=layout.circle, main = "Output directed graph")
par(old.par)
```

---

pairwiseMatrix	<i>Pairwise plotting of multivariate data</i>
----------------	-----------------------------------------------

---

## Description

Display a pairwise scatter plot of two datasets for a random selection of variables. If the second dataset is not given, the function displays a histogram with normal curve superposition.

## Usage

```
pairwiseMatrix(x, y = NULL, size = nrow(x), r = 4, c = 4, ...)
```

## Arguments

x	A matrix or data.frame (n x p) of continuous data.
y	A matrix or data.frame (n x q) of continuous data.
size	number of rows to be sampled (default s = nrow(z)).
r	number of rows of the plot layout (default r = 4).
c	number of columns of the plot layout (default r = 4).
...	Currently ignored.

## Value

No return value

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## Examples

```
adjdata <- SEMbap(sachs$graph, log(sachs$pkc))$data
rawdata <- log(sachs$pkc)
pairwiseMatrix(adjdata, rawdata, size = 1000)
```

---

parameterEstimates	<i>Parameter Estimates of a fitted SEM</i>
--------------------	--------------------------------------------

---

### Description

Wrapper of the lavaan parameterEstimates() function for RICF and CGGM algorithms

### Usage

```
parameterEstimates(fit, ...)
```

### Arguments

fit	A RICF or constrained GGM fitted model object.
...	Currently ignored.

### Value

A data.frame containing the estimated parameters

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

### Examples

```
ricf1 <- SEMrun(sachs$graph, log(sachs$pkc), sachs$group, algo = "ricf")
parameterEstimates(ricf1$fit)

cggm1 <- SEMrun(sachs$graph, log(sachs$pkc), sachs$group, algo = "cggm")
parameterEstimates(cggm1$fit)
```

---

pathFinder	<i>Perturbed path search utility</i>
------------	--------------------------------------

---

### Description

This function uses [SEMace](#) to find significant causal effects between source-sink pairs and [SEMpath](#) to fit them and test their edge perturbation.



**Usage**

```
pathFinder(
  graph,
  data,
  group = NULL,
  ace = NULL,
  path = "directed",
  method = "BH",
  alpha = 0.05,
  ...
)
```

**Arguments**

graph	Input network as an igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
group	group A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. Group specification enables edge perturbation testing. By default, group = NULL.
ace	A data.frame generated by <a href="#">SEMace</a> . If NULL, <a href="#">SEMace</a> will be automatically run.
path	If path = "directed", all directed paths between the two nodes will be included in the fitted model. If path = "shortest", only shortest paths will be considered.
method	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method = "BH" (i.e., FDR multiple test correction).
alpha	Significance level for ACE selection (by default, alpha = 0.05).
...	Currently ignored.

**Value**

A list of 3 objects:

- "paths", list of paths as igraph objects;
- "fit", fitting results for each path as a lavaan object;
- "dfp", a data.frame containing SEM global fitting statistics.

**Author(s)**

Fernando Palluzzi <fernando.palluzzi@gmail.com>

**Examples**

```
# Find and evaluate significantly perturbed paths
```

```

library(huge)
als.npn <- huge.npn(alsData$exprs)

adjData <- SEMbap(graph = alsData$graph, data = als.npn)$data

ace <- SEMace(graph = alsData$graph, data = adjData,
             group = alsData$group)
ace <- ace[order(ace$pvalue),]
print(ace)

paths <- pathFinder(graph = alsData$graph, data = adjData,
                   group = alsData$group,
                   ace = ace)

print(paths$dfp)
head(parameterEstimates(paths$fit[[1]]))
gplot(paths$paths[[1]])

```

---

properties

*Graph properties summary and graph decomposition*


---

### Description

Produces a summary of network properties and returns graph components (ordered by decreasing size), without self-loops.

### Usage

```
properties(graph, data = NULL, ...)
```

### Arguments

graph	Input network as an igraph object.
data	An optional data matrix (default data = NULL) with rows corresponding to subjects, and columns to graph nodes (variables). Nodes will be mapped onto variable names.
...	Currently ignored.

### Value

List of graph components, ordered by decreasing size (the first component is the giant one), without self-loops.

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

## Examples

```
# Extract the "Type II diabetes mellitus" pathway:
g <- kegg.pathways[["Type II diabetes mellitus"]]
summary(g)
properties(g)
```

---

resizeGraph

*Interactome-assisted graph re-seizing*


---

## Description

An input directed graph is re-sized, removing edges or adding edges/nodes. This function takes three input graphs: the first is the input causal model (i.e., a directed graph), and the second can be either a directed or undirected graph, providing a set of connections to be checked against a directed reference network (i.e., the third input) and imported to the first graph.

## Usage

```
resizeGraph(g = list(), gnet, d = 2, v = TRUE, verbose = FALSE, ...)
```

## Arguments

<code>g</code>	A list of two graphs as igraph objects, <code>g=list(graph1, graph2)</code> .
<code>gnet</code>	External directed network as an igraph object. The reference network should have weighted edges, corresponding to their interaction p-values, as an edge attribute <code>E(gnet)\$pv</code> . Then, connections in <code>graph2</code> will be checked by known connections from the reference network, intercepted by the minimum-weighted shortest path found among the equivalent ones by the Dijkstra algorithm, as implemented in the <b>igraph</b> function <code>all_shortest_paths()</code> .
<code>d</code>	An integer value indicating the maximum geodesic distance between two nodes in the interactome to consider the inferred interaction between the same two nodes in <code>graph2</code> as validated, otherwise the edges are removed. For instance, if <code>d = 2</code> , two interacting nodes must either share a direct interaction or being connected through at most one mediator in the reference interactome (in general, at most <code>d - 1</code> mediators are allowed). Typical <code>d</code> values include 2 (at most one mediator), or <code>mean_distance(gnet)</code> (i.e., the average shortest path length for the reference network). Setting <code>d = 0</code> , is equivalent to <code>gnet = NULL</code> .
<code>v</code>	A logical value. If <code>TRUE</code> (default) new nodes and edges on the validated shortest path in the reference interactome will be added in the re-sized graph.
<code>verbose</code>	A logical value. If <code>FALSE</code> (default), the processed graphs will not be plotted to screen, saving execution time (for large graphs)
<code>...</code>	Currently ignored.

## Details

Typically, the first graph is an estimated causal graph (DAG), and the second graph is the output of `SEMdag` or an external covariance graph. In the former we use the new inferred causal structure stored in the `dag.new` object. In the latter, we use the new inferred covariance structure stored in the covariance graph object. Both directed (causal) edges and covariances (i.e., bidirected edges) highlight emergent hidden topological proprieties, absent in the input graph. Estimated directed edges between nodes *X* and *Y* are interpreted as either direct links or direct paths mediated by hidden connector nodes. Covariances between any two bow-free nodes *X* and *Y* may hide causal relationships, not explicitly represented in the current model. Conversely, directed (or bi-directed) edges could be redundant or artifact, specific to the observed data and could be deleted. Function `resizeGraph()` leverage on these concepts to extend/reduce a causal model, importing new connectors or deleting estimated edges, if they are present or absent in a given reference network. The whole process may lead to the discovery of new paths of information flow, and cut edges not corroborate by a validated network. Since added nodes can already be present in the causal graph, network resize may create cross-connections between old and new paths and their possible closure into circuits.

## Value

"Ug", the re-sized graph, the graph union of the causal graph `graph1` and the re-sized graph `graph2`

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38 (20), 4829–4830 <<https://doi.org/10.1093/bioinformatics/btac567>>

## Examples

```
# Extract the "Protein processing in endoplasmic reticulum" pathway:

g <- kegg.pathways[["Protein processing in endoplasmic reticulum"]]
G <- properties(g)[[1]]; summary(G)

# Extend a graph using new inferred DAG edges (dag+dag.new):

library(huge)
als.npn <- huge.npn(alsData$exprs)

dag <- SEMdag(graph = G, data = als.npn, beta = 0.1)
gplot(dag$dag)
ext <- resizeGraph(g=list(dag$dag, dag$dag.new), gnet = kegg, d = 2)
gplot(ext)

# Create a directed graph from correlation matrix, using
```

```

# i) an empty graph as causal graph,
# ii) a covariance graph,
# iii) KEGG as reference:

corr2graph<- function(R, n, alpha=5e-6, ...)
{
  Z <- qnorm(alpha/2, lower.tail=FALSE)
  thr <- (exp(2*Z/sqrt(n-3))-1)/(exp(2*Z/sqrt(n-3))+1)
  A <- ifelse(abs(R) > thr, 1, 0)
  diag(A) <- 0
  return(graph_from_adjacency_matrix(A, mode="undirected"))
}

v <- which(colnames(als.npn) %in% V(G)$name)
selectedData <- als.npn[, v]
G0 <- make_empty_graph(n = ncol(selectedData))
V(G0)$name <- colnames(selectedData)
G1 <- corr2graph(R = cor(selectedData), n= nrow(selectedData))
ext <- resizeGraph(g=list(G0, G1), gnet = kegg, d = 2, v = TRUE)

#Graphs
old.par <- par(no.readonly = TRUE)
par(mfrow=c(1,2), mar=rep(1,4))
plot(G1, layout = layout.circle)
plot(ext, layout = layout.circle)
par(old.par)

```

---

sachs

*Sachs multiparameter flow cytometry data and consensus model*


---

## Description

Flow cytometry data and causal model from Sachs et al. (2005).

## Usage

```
sachs
```

## Format

"sachs" is a list of 5 objects:

1. "rawdata", a list of 14 data.frames containing raw flow cytometry data (Sachs et al., 2005);
2. "graph", consensus signaling network;
3. "model", consensus model (lavaan syntax);

4. "pkc", data.frame of 1766 samples and 11 variables, containing cd3cd28 (baseline) and pma (PKC activation) data;
5. "group", a binary group vector, where 0 is for cd3cd28 samples (n = 853) and 1 is for pma samples (n = 913).
6. "details", a data.frame containing dataset information.

## Source

[doi:10.1126/science.1105809](https://doi.org/10.1126/science.1105809)

## References

Sachs K, Perez O, Pe'er D, Lauffenburger DA, Nolan GP (2019). Causal Protein-Signaling Networks Derived from Multiparameter Single-Cell Data. *Science*, 308(5721): 523-529.

## Examples

```
# Dataset content
names(sachs$rawdata)
dim(sachs$pkc)
table(sachs$group)
cat(sachs$model)
gplot(sachs$graph)
```

---

sem.mediation

*SEM-based mediation analysis*

---

## Description

Perform mediation analysis through structural equation modeling (SEM). The SEM is represented by two equations:  $x_2 \sim b \cdot x_1$  and  $y \sim a \cdot x_1 + c \cdot x_2$ , where  $x_1$  is the exposure factor,  $y$  is the outcome variable, and  $x_2$  is the mediator. This model is iteratively applied for each column in the input data matrix, given  $x_1$  and  $y$ . A new parameter  $IE := b \cdot c$  is defined, corresponding to the mediation effect.

## Usage

```
sem.mediation(
  data,
  x1,
  y,
  model = NULL,
  group = NULL,
  fixed.x = TRUE,
  se = "robust.huber.white",
  std.x = FALSE,
  std.y = FALSE,
  std.m = FALSE,
```

```

    adjust.method = "BH",
    gene.symbols = FALSE,
    ...
)

```

## Arguments

<code>data</code>	A matrix with rows corresponding to subjects, and columns to potential mediators.
<code>x1</code>	A numeric vector corresponding to the exposure factor.
<code>y</code>	A numeric vector corresponding to the outcome variable.
<code>model</code>	The user might specify a different model. To use the given mediation model, leave this argument to NULL (default).
<code>group</code>	A vector to group subjects for multigroup analysis. By default, <code>group = NULL</code> .
<code>fixed.x</code>	Logical value. If TRUE (default), the exposure factor (i.e., the exogenous variable) estimates are fixed to its sampling values. If FALSE, their estimates are considered as free parameters.
<code>se</code>	Standard error estimation method. Use "standard" (default) for conventional standard errors, based on the inverse information matrix. Use "robust.huber.white" for pseudo ML Huber-White estimation.
<code>std.x</code>	Logical value. If TRUE, the exposure factor is standardized (default = FALSE).
<code>std.y</code>	Logical value. If TRUE, the outcome variable is standardized (default = FALSE).
<code>std.m</code>	Logical value. If TRUE, the mediator is standardized (default = FALSE).
<code>adjust.method</code>	One of the multiple testing correction methods from the <code>p.adjust</code> function. By default, the Benjamini-Hochberg ("BH") adjustment method is used.
<code>gene.symbols</code>	Logical value. If variables are genes and data matrix column names are entrez gene IDs, this argument can be set to TRUE to obtain a new column with official gene symbols (default = FALSE).
<code>...</code>	Currently ignored.

## Value

A list of two objects:

1. "model", The output model as an `igraph` object;
2. "clusters", A list containing model clustering results. It includes three more objects:
  - "membership", A vector reporting cluster membership for each node;
  - "fit", hidden model fitting as a `lavaan` object;
  - "dataHM", data matrix containing cluster scores (hidden variables) alongside the original data variables.

## Author(s)

Fernando Palluzzi <fernando.palluzzi@gmail.com>

## Examples

```
library(huge)
library(org.Hs.eg.db)

als.npn <- huge.npn(alsData$exprs)

# Random sampling 10 variables (possible mediators)
j <- sample.int(ncol(als.npn), 10)

# Generate a random binary exposure vector
exposure <- rbinom(n = nrow(als.npn), size = 1, prob = 0.3)

# Generate a random gaussian outcome vector
outcome <- rnorm(n = nrow(als.npn))

mediators <- sem.mediation(data = als.npn[, j],
                           x1 = exposure,
                           y = outcome,
                           gene.symbols = TRUE)
```

---

SEMace

---

*Compute the Average Causal Effect (ACE) for a given source-sink pair*


---

## Description

Compute total effects as ACEs of variables X on variables Y in a directed acyclic graph (DAG). The ACE will be estimated as the path coefficient of X (i.e.,  $\theta$ ) in the linear equation  $Y \sim X + Z$ . The set Z is defined as the adjustment (or conditioning) set of Y over X, applying various adjustment sets. Standard errors (SE), for each ACE, are computed following the `lm` standard procedure or a bootstrap-based procedure (see [boot](#) for details).

## Usage

```
SEMace(
  graph,
  data,
  group = NULL,
  type = "parents",
  effect = "all",
  method = "BH",
  alpha = 0.05,
  boot = NULL,
  ...
)
```



**Arguments**

graph	An igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. If group = NULL (default), group influence will not be considered.
type	character Conditioning set Z. If "parents" (default) the Pearl's back-door set (Pearl, 1998), "minimal" the dagitty minimal set (Perkovic et al, 2018), or "optimal" the O-set with the smallest asymptotic variance (Witte et al, 2020) are computed.
effect	character X to Y effect. If "all" (default) all effects from X to Y, "source2sink" only effects from source X to sink Y, or "direct" only direct effects from X to Y are computed.
method	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method = "BH" (i.e., FDR multiple test correction).
alpha	Significance level for ACE selection (by default, alpha = 0.05).
boot	The number of bootstrap samplings enabling bootstrap computation of ACE standard errors. If NULL (default), bootstrap is disabled.
...	Currently ignored.

**Value**

A data.frame of ACE estimates between network sources and sinks.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

- Pearl J (1998). Graphs, Causality, and Structural Equation Models. *Sociological Methods & Research*, 27(2):226-284. <<https://doi.org/10.1177/0049124198027002004>>
- Perkovic E, Textor J, Kalisch M, Maathuis MH (2018). Complete graphical characterization and construction of adjustment sets in Markov equivalence classes of ancestral graphs. *Journal of Machine Learning Research*, 18:1-62. <<http://jmlr.org/papers/v18/16-319.html>>
- Witte J, Henckel L, Maathuis MH, Didelez V (2020). On efficient adjustment in causal graphs. *Journal of Machine Learning Research*, 21:1-45. <<http://jmlr.org/papers/v21/20-175.htm>>

**Examples**

```
# ACE without group, O-set, all effects:
ace1 <- SEMace(graph = sachs$graph, data = log(sachs$pkc),
               group = NULL, type = "optimal", effect = "all",
               method = "BH", alpha = 0.05, boot = NULL)
print(ace1)
```

```
# ACE with group perturbation, Pa-set, direct effects:
ace2 <- SEMace(graph = sachs$graph, data = log(sachs$pkc),
               group = sachs$group, type = "parents", effect = "direct",
               method = "none", alpha = 0.05, boot = NULL)
print(ace2)
```

SEMbap

*Bow-free covariance search and data de-correlation*

## Description

SEMbap() function implements different deconfounding methods to adjust the data matrix by removing latent sources of confounding encoded in them. The selected methods are either based on: (i) Bow-free Acyclic Paths (BAP) search, (ii) LVs proxies as additional source nodes of the data matrix, Y or (iii) spectral transformation of Y.

## Usage

```
SEMbap(
  graph,
  data,
  group = NULL,
  dalgo = "cggm_dsep",
  method = "BH",
  alpha = 0.05,
  hcount = "auto",
  limit = 30000,
  verbose = FALSE,
  ...
)
```

## Arguments

graph	An igraph object.
data	A matrix with rows corresponding to subjects, and or <a href="#">SILGGM</a> columns to graph nodes (variables).
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. If NULL (default), confounding within group will not be considered.
dalgo	Deconfounding method. Six algorithms are available: <ul style="list-style-type: none"> <li>• "cggm_dsep" (default). The algorithm make: (i) exhaustive search of bow-free significant covariances (see details) through <a href="#">Shipley.test</a> function; (ii) estimation of the inverse of the selected covariance matrix (i.e. the precision matrix, W) through <a href="#">fitConGraph</a> function; (iii) obtain the de-correlated data matrix, Z by multiplying the data matrix, Y rightward by the</li> </ul>

square root of the estimated precision matrix,  $Z=YW^{(1/2)}$  as suggested by Grassi, Palluzzi and Tarantino (2022).

- "glpc\_dsep". The algorithm first makes an exhaustive search of bow-free significant covariances through [ShipleY.test](#) function. Once obtained the adjacency matrix, Graph-Laplacian PCA (gLPCA) algorithm learns a low dimensional representation of the observed data matrix that incorporates graph structure (Jiang et al., 2013). Then, the DAG is extended by including the confounding proxies, i.e. LVs, as additional source nodes defined by last  $q$  principal component scores of gLPCA and these LV scores are added to the data matrix,  $Z=cbind(LV,Y)$ .
- "pc". The procedure add additional source nodes to DAG as in "glpc\*" algorithms, but confounding proxies are the  $q$  principal component scores extracted by Spectral decomposition (SVD) selecting only graph nodes and without graph edge information and bow-free covariance search.
- "trim" or "pcss". Cévid et al. (2020) suggest multiplying the data matrix,  $Y$  leftward by a well selected spectrum transformation matrix,  $T$  which modifies the singular values of  $Y$ , while keeping its singular vectors intact,  $Z=TY$ . Trim transform limits all singular values to be at most some constant ( $t$ ), where  $t$  = median of the singular values. While, PCSS transform set equal to zero the first  $q$  singular values, and keep the others intact.

method	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method is set to "BH" (i.e., Benjamini-Hochberg multiple test correction).
alpha	Significance level for false discovery rate (FDR) used for d-separation or CI tests. This argument is used to control data de-correlation. A higher alpha level includes more hidden covariances, thus considering more sources of confounding. If $\alpha = 0$ , data de-correlation is disabled. By default, $\alpha = 0.05$ .
hcount	The number of latent (or hidden) variables. By default <code>hcount="auto"</code> , the hidden count is determined with a permutation method (see details). Currently ignored if ( <code>dalgo="cggm"</code> or <code>"trim"</code> ).
limit	An integer value corresponding to the number of missing edges tolerance. Beyond this limit, multicore computation is enabled to reduce the computational burden of the exhaustive BAP search of the <a href="#">ShipleY.test</a> procedure. By default, <code>limit = 30000</code> .
verbose	A logical value. If FALSE (default), the processed graphs will not be plotted to screen.
...	Currently ignored.

## Details

Missing edges in causal network inference using a directed acyclic graph (DAG) are frequently hidden by unmeasured confounding variables. A Bow-free Acyclic Paths (BAP) search is performed with d-separation or Conditional Independence (CI) tests between all pairs of variables with missing connection in the input DAG, adding a bidirected edge (i.e., bow-free covariance) to the DAG when there is an association between them. The d-separation test evaluates if two variables ( $Y1$ ,  $Y2$ ) in a DAG are conditionally independent for a given conditioning set,  $C$  represented in a DAG by the union of the parent sets of  $Y1$  and  $Y2$  (ShipleY, 2000). The CI test (Jankova and van de Geer,

2015) evaluates if two variables (Y1, Y2) in an undirected graph are independent given the set = "rest" (the other variables excluding Y1 and Y2). A new bow-free covariance is added if there is a significant (Y1, Y2) association at a significance level  $\alpha$ , after multiple testing correction. The selected covariance between pairs of nodes is interpreted as the effect of a latent variable (LV) acting on both nodes; i.e., the LV is an unobserved confounder. BAP-based algorithms adjust (or de-correlate) the observed data matrix by conditioning out the latent triggers responsible for the nuisance edges. For "pc" algorithm the number of hidden proxies,  $q$  is determined by a permutation method. It compares the singular values to what they would be if the variables were independent, which is estimated by permuting the columns of the data matrix,  $Y$  and selects components if their singular values are larger than those of the permuted data (for a review see Dobriban, 2020). While for "glpc\*" algorithms,  $q$  is determined by the number of clusters by spectral clustering through `cluster_leading_eigen` function. If the input graph is not acyclic, a warning message will be raised, and a cycle-breaking algorithm will be applied (see `graph2dag` for details).

### Value

A list of four objects:

- "dag", the directed acyclic graph (DAG) extracted from input graph. If (dalgo = "glpc\*" or "pc"), the DAG also includes LVs as source nodes.
- "dsep", the data.frame of all d-separation tests or the conditional independence (CI) tests over missing edges in the DAG. If (dalgo = "pc" or "trim"), dsep dataframe is equal to NULL.
- "adj", the adjacency matrix of selected bow-free covariances; i.e, the missing edges selected after multiple testing correction. If (dalgo = "pc" or "trim"), adj matrix is equal to NULL.
- "data", the adjusted (de-correlated) data matrix or if (dalgo = "glpc\*", or "pc"), the combined data matrix, where the first columns represent LVs scores and the other columns are the raw data.

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

### References

Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38(20), 4829–4830. <<https://doi.org/10.1093/bioinformatics/btac567>>

Shipley B (2000). A new inferential test for path models based on DAGs. *Structural Equation Modeling*, 7(2), 206-218. <[https://doi.org/10.1207/S15328007SEM0702\\_4](https://doi.org/10.1207/S15328007SEM0702_4)>

Jankova J, van de Geer S. (2015) Confidence intervals for high-dimensional inverse covariance estimation. *Electronic Journal of Statistics*, 9, 1205-1229. <<https://doi.org/10.1214/15-EJS1031>>

Jiang B, Ding C, Bin L, Tang J (2013). Graph-Laplacian PCA: Closed-Form Solution and Robustness. *IEEE Conference on Computer Vision and Pattern Recognition*, 3492-3498. <<https://doi.org/10.1109/CVPR.2013.448>>

Ćevic D, Bühlmann P, Meinshausen N (2020). Spectral deconfounding via perturbed sparse linear models. *J. Mach. Learn. Res*, 21 (232), 1-41. <<http://jmlr.org/papers/v21/19-545.html>>

Dobriban E (2020). Permutation methods for Factor Analysis and PCA. *Ann. Statist.* 48(5): 2824-2847 <<https://doi.org/10.1214/19-AOS1907>>

**Examples**

```
#Set parameters
graph <- sachs$graph
data <- log(sachs$pkc)
group <-sachs$group

# BAP search with CGGM and d-separation test (default)
bap <- SEMbap(graph, data, verbose = TRUE)
#gplot(bap$dag)

# BAP search with gLPCA scores and CI test
glpc <- SEMbap(graph, data, dalgo = "glpc_ci")
gplot(glpc$dag)

# Model fitting (with node-perturbation)
sem1 <- SEMrun(graph, data, group)
bap1 <- SEMrun(bap$dag, bap$data, group)
glpc1 <- SEMrun(glpc$dag, glpc$data, group)
```

SEMdag

*Estimate the optimal DAG from an input graph***Description**

Extract the optimal DAG from an input graph, using topological order or top-down order search and LASSO-based algorithm, implemented in [glmnet](#).

**Usage**

```
SEMdag(
  graph,
  data,
  LO = "TO",
  beta = 0,
  lambdas = NA,
  penalty = TRUE,
  verbose = FALSE,
  ...
)
```

**Arguments**

graph	An igraph object.
data	A matrix with n rows corresponding to subjects, and p columns to graph nodes (variables).

LO	character for linear order method. If LO="TO" the topological order of the input DAG is enabled (default), while LO="TD" the data-driven top-down minimum conditional variance method is performed.
beta	Numeric value. Minimum absolute LASSO beta coefficient for a new interaction to be retained in the final model. By default, beta = 0.
lambdas	A vector of regularization LASSO lambda values. If lambdas is NULL, the <a href="#">glmnet</a> default using cross-validation lambdas is enabled. If lambdas is NA (default), the tuning-free scheme is enabled by fixing lambdas = sqrt(log(p)/n), as suggested by Janková and van de Geer (2015) and many others. This will both reduce computational time and provide the same result at each run.
penalty	A logical value. Separate penalty factors can be applied to each coefficient. This is a number that multiplies lambda to allow differential shrinkage. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. If TRUE (default) weights are based on the graph edges: 0 (i.e., edge present) and 1 (i.e., missing edge) ensures that the input edges will be retained in the final model. If FALSE the <a href="#">glmnet</a> default is enabled (all weights equal to 1). Note: the penalty factors are internally rescaled to sum p (the number of variables).
verbose	A logical value. If FALSE (default), the processed graphs will not be plotted to screen.
...	Currently ignored.

### Details

The optimal DAG is estimated using the order search approach. First a linear order of  $p$  nodes is determined, and from this sort, the DAG can be learned using successive penalized (L1) regressions (Shojaie and Michailidis, 2010). The estimate linear order are obtained from *a priori* graph topological order (TO), or with a data-driven high dimensional top-down (TD) approach (best subset regression), assuming a SEM whose error terms have equal variances (Peters and Bühlmann, 2014; Chen et al, 2019). If the input graph is not acyclic, a warning message will be raised, and a cycle-breaking algorithm will be applied (see [graph2dag](#) for details). Output DAG edges will be colored in gray, if they were present in the input graph, and in green, if they are new edges generated by LASSO screening.

### Value

A list of 3 igraph objects:

1. "dag", the estimated DAG;
2. "dag.new", new estimated connections;
3. "dag.old", connections preserved from the input graph.

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

- Tibshirani R, Bien J, Friedman J, Hastie T, Simon N, Taylor J, Tibshirani RJ (2012). Strong rules for discarding predictors in lasso type problems. *Royal Statistical Society: Series B (Statistical Methodology)*, 74(2): 245-266. <<https://doi.org/10.1111/j.1467-9868.2011.01004.x>>
- Shojaie A, Michailidis G (2010). Penalized likelihood methods for estimation of sparse high-dimensional directed acyclic graphs. *Biometrika*, 97(3): 519-538. <<https://doi.org/10.1093/biomet/asq038>>
- Jankova J, van de Geer S (2015). Confidence intervals for high-dimensional inverse covariance estimation. *Electronic Journal of Statistics*, 9(1): 1205-1229. <<https://doi.org/10.1214/15-EJS1031>>
- Peters J, Bühlmann P (2014). Identifiability of Gaussian structural equation models with equal error variances. *Biometrika*, 101(1):219–228.
- Chen W, Drton M, Wang YS (2019). On Causal Discovery with an Equal-Variance Assumption. *Biometrika*, 106(4): 973-980.

## See Also

[modelSearch](#)

## Examples

```
# DAG estimation
G <- SEMdag(graph = sachs$graph, data = log(sachs$pkc), beta = 0.05)

# Model fitting
sem <- SEMrun(graph = G$dag, data = log(sachs$pkc), group = sachs$group)

# Graphs
old.par <- par(no.readonly = TRUE)
par(mfrow=c(2,2), mar=rep(1,4))
plot(sachs$graph, layout=layout.circle, main="input graph")
plot(G$dag, layout=layout.circle, main = "Output DAG")
plot(G$dag.old, layout=layout.circle, main = "Inferred old edges")
plot(G$dag.new, layout=layout.circle, main = "Inferred new edges")
par(old.par)
```

## Description

Creates a network with perturbed edges obtained from the output of [SEMace](#), comparable to the procedure in Jablonski et al (2022), or of [SEMrun](#) with two-group and CGGM solver, comparable to the algorithm 2 in Belyaeva et al (2021). To increase the efficiency of computations for large graphs, users can select to break the network structure into clusters, and select the topological clustering method (see [clusterGraph](#)). The function [SEMrun](#) is applied iteratively on each cluster (with size min > 10 and max < 500) to obtain the graph with the full list of perturbed edges.

**Usage**

```
SEMdci(graph, data, group, type = "ace", method = "BH", alpha = 0.05, ...)
```

**Arguments**

<code>graph</code>	Input network as an igraph object.
<code>data</code>	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
<code>group</code>	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects.
<code>type</code>	Average Causal Effect (ACE) with two-group, "parents" (back-door) adjustment set, and "direct" effects ( <code>type = "ace"</code> , default), or CGGM solver with two-group using a clustering method. If <code>type = "tahc"</code> , network modules are generated using the tree agglomerative hierarchical clustering method. Other non-tree clustering methods from igraph package include: "wtc" (walktrap community structure with short random walks), "ebc" (edge betweenness clustering), "fgc" (fast greedy method), "lbc" (label propagation method), "lec" (leading eigenvector method), "loc" (multi-level optimization), "opc" (optimal community structure), "sgc" (springlass statistical mechanics), "none" (no breaking network structure into clusters).
<code>method</code>	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method is set to "BH" (i.e., FDR multiple test correction).
<code>alpha</code>	Significance level (default = 0.05) for edge set selection.
<code>...</code>	Currently ignored.

**Value**

An igraph object.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

Belyaeva A, Squires C, Uhler C (2021). DCI: learning causal differences between gene regulatory networks. *Bioinformatics*, 37(18): 3067–3069. <<https://doi.org/10.1093/bioinformatics/btab167>>

Jablonski K, Pirkl M, Čevič D, Bühlmann P, Beerenwinkel N (2022). Identifying cancer pathway dysregulations using differential causal effects. *Bioinformatics*, 38(6):1550–1559. <<https://doi.org/10.1093/bioinformatics/btad167>>

**Examples**

```
## Not run:

# Nonparanormal(npn) transformation
library(huge)
als.npn <- huge.npn(alsData$exprs)
```



```

g <- kegg.pathways[["MAPK signaling pathway"]]
G <- properties(g)[[1]]; summary(G)

# Create ALS network with perturbed edges using edge betweenness clustering
gU<- SEMdci(G, als.npn, alsData$group, type="ebc", method="BH", alpha=0.2)
gcU<- properties(gU)

old.par <- par(no.readonly = TRUE)
par(mfrow=c(2,2), mar=rep(1,4))
gplot(gcU[[1]], l="fdp") # max component
gplot(gcU[[2]], l="fdp") # 2nd component
gplot(gcU[[3]], l="fdp") # 3rd component
gplot(gcU[[4]], l="fdp") # 4th component
par(old.par)

## End(Not run)

```

SEMgsa

*SEM-based gene set analysis (GSA)*

## Description

Gene Set Analysis (GSA) via self-contained test for group effect on signaling (directed) pathways based on SEM. The core of the methodology is implemented in the RICF algorithm of `SEMrun()`, recovering from RICF output node-specific group effect p-values, and Brown's combined permutation p-values of node activation and inhibition.

## Usage

```
SEMgsa(g = list(), data, group, method = "BH", alpha = 0.05, n_rep = 1000, ...)
```

## Arguments

<code>g</code>	A list of pathways to be tested.
<code>data</code>	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
<code>group</code>	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects.
<code>method</code>	Multiple testing correction method. One of the values available in <a href="#">p.adjust</a> . By default, method is set to "BH" (i.e., Benjamini-Hochberg correction).
<code>alpha</code>	Gene set test significance level (default = 0.05).
<code>n_rep</code>	Number of randomization replicates (default = 1000).
<code>...</code>	Currently ignored.

## Details

For gaining more biological insights into the functional roles of pre-defined subsets of genes, node perturbation obtained from RICF fitting has been combined with up- or down-regulation of genes from KEGG to obtain overall pathway perturbation as follows:

- The node perturbation is defined as activated when the minimum among the p-values is positive; if negative, the status is inhibited.
- Up- or down- regulation of genes (derived from KEGG database) has been obtained from the weighted adjacency matrix of each pathway as column sum of weights over each source node. If the overall sum of node weights is below 1, the pathway is flagged as down-regulated otherwise as up-regulated.
- The combination between these two quantities allows to define the direction (up or down) of gene perturbation. Up- or down regulated gene status, associated with node inhibition, indicates a decrease in activation (or increase in inhibition) in cases with respect to control group. Conversely, up- or down regulated gene status, associated with node activation, indicates an increase in activation (or decrease in inhibition) in cases with respect to control group.

## Value

A list of 2 objects:

1. "gsa", A data.frame reporting the following information for each pathway in the input list:
  - "No.nodes", pathway size (number of nodes);
  - "No.DEGs", number of differential expression genes (DEGs) within the pathway, after multiple test correction with one of the methods available in [p.adjust](#);
  - "pert", pathway perturbation status (see details);
  - "pNA", Brown's combined P-value of pathway node activation;
  - "pNI", Brown's combined P-value of pathway node inhibition;
  - "PVAL", Bonferroni combined P-value of pNA, and pNI; i.e.,  $2 * \min(pNA, PNI)$ ;
  - "ADJP", Adjusted Bonferroni P-value of pathway perturbation; i.e.,  $\min(\text{No.pathways} * PVAL; 1)$ .
2. "DEG", a list with DEGs names per pathways.

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

Grassi, M., Tarantino, B. SEMgsa: topology-based pathway enrichment analysis with structural equation models. BMC Bioinformatics 23, 344 (2022). <https://doi.org/10.1186/s12859-022-04884-8>

**Examples**

```
## Not run:

# Nonparanormal(npn) transformation

library(huge)
als.npn <- huge.npn(alsData$exprs)

# Selection of FTD-ALS pathways from kegg.pathways.Rdata

paths.name <- c("MAPK signaling pathway",
                "Protein processing in endoplasmic reticulum",
                "Endocytosis",
                "Wnt signaling pathway",
                "Neurotrophin signaling pathway",
                "Amyotrophic lateral sclerosis")

j <- which(names(kegg.pathways) %in% paths.name)

GSA <- SEMgsa(kegg.pathways[j], als.npn, alsData$group,
              method = "bonferroni", alpha = 0.05,
              n_rep = 1000)

GSA$gsa
GSA$DEG

## End(Not run)
```

---

SEMPath	<i>Search for directed or shortest paths between pairs of source-sink nodes</i>
---------	---------------------------------------------------------------------------------

---

**Description**

Find and fit all directed or shortest paths between two source-sink nodes of a graph.

**Usage**

```
SEMPath(graph, data, group, from, to, path, verbose = FALSE, ...)
```

**Arguments**

graph	An igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
group	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. If NULL (default), group influence will not be considered.

from	Starting node name (i.e., source node).
to	Ending node name (i.e., sink node).
path	If path = "directed", all directed paths between the two nodes will be included in the fitted model. If path = "shortest", only shortest paths will be returned.
verbose	Show the directed (or shortest) path between the given source-sink pair inside the input graph.
...	Currently ignored.

### Value

A list of four objects: a fitted model object of class `lavaan` ("fit"), aggregated and node-specific group effect estimates and P-values ("gest"), the extracted subnetwork as an `igraph` object ("graph"), and the input graph with a color attribute mapping the chosen path ("map").

### Author(s)

Mario Grassi <mario.grassi@unipv.it>

### Examples

```
# Directed path fitting
path <- SEMpath(graph = sachs$graph, data = log(sachs$pkc),
               group = sachs$group,
               from = "PIP3",
               to = "Erk",
               path = "directed")

# Summaries
summary(path$fit)
print(path$gest)

# Graphs
gplot(path$map, main="path from PiP2 to Erk")
plot(path$map, layout=layout.circle, main="path from PiP2 to Erk")
```

### Description

SEMrun() converts a (directed, undirected, or mixed) graph to a SEM and fits it. If a binary group variable (i.e., case/control) is present, node-level or edge-level perturbation is evaluated. This function can handle loop-containing models, although multiple links between the same two nodes (including self-loops and mutual interactions) and bows (i.e., a directed and a bidirected link between two nodes) are not allowed.

**Usage**

```
SEMrun(
  graph,
  data,
  group = NULL,
  fit = 0,
  algo = "lavaan",
  start = NULL,
  SE = "standard",
  n_rep = 1000,
  limit = 100,
  ...
)
```

**Arguments**

<code>graph</code>	An igraph object.
<code>data</code>	A matrix with rows corresponding to subjects, and columns to graph nodes (variables).
<code>group</code>	A binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. If <code>NULL</code> (default), group influence will not be considered.
<code>fit</code>	A numeric value indicating the SEM fitting mode. If <code>fit = 0</code> (default), no group effect is considered. If <code>fit = 1</code> , a "common" model is used to evaluate group effects on graph nodes. If <code>fit = 2</code> , a two-group model is used to evaluate group effects on graph edges.
<code>algo</code>	MLE method used for SEM fitting. If <code>algo = "lavaan"</code> (default), the SEM will be fitted using the NLMINB solver from lavaan R package, with standard errors derived from the expected Fisher information matrix. If <code>algo = "ricf"</code> , the model is fitted via residual iterative conditional fitting (RICF; Drton et al. 2009). If <code>algo = "cggm"</code> , model fitting is based on constrained Gaussian Graphical Modeling (CGGM; Hastie et al. 2009, p. 446).
<code>start</code>	Starting value of SEM parameters for <code>algo = "lavaan"</code> . If <code>start</code> is <code>NULL</code> (default), the algorithm will determine the starting values. If <code>start</code> is a numeric value, it will be used as a scaling factor for the edge weights in the graph object (graph attribute <code>E(graph)\$weight</code> ). For instance, a scaling factor is useful when weights have fixed values (e.g., 1 for activated, -1 for repressed, and 0 for unchanged interaction). Fixed values may compromise model fitting, and scaling them is a safe option to avoid this problem. As a rule of thumb, to our experience, <code>start = 0.1</code> generally performs well with -1, 0, 1 weights.
<code>SE</code>	If "standard" (default), with <code>algo = "lavaan"</code> , conventional standard errors are computed based on inverting the observed information matrix. If "none", no standard errors are computed.
<code>n_rep</code>	Number of randomization replicates (default = 1000), for permutation flip or bootstrap samples, if <code>algo = "ricf"</code> .

limit	An integer value corresponding to the network size (i.e., number of nodes). Beyond this limit, the execution under <code>algo = "lavaan"</code> will run with <code>SE = "none"</code> , if <code>fit = 0</code> , or will be redirected to <code>algo = "ricf"</code> , if <code>fit = 1</code> , or to <code>algo = "cggm"</code> , if <code>fit = 2</code> . This redirection is necessary to reduce the computational demand of standard error estimation by lavaan. Increasing this number will enforce lavaan execution when <code>algo = "lavaan"</code> .
...	Currently ignored.

## Details

SEMrun maps data onto the input graph and converts it into a SEM. Directed connections ( $X \rightarrow Y$ ) are interpreted as direct causal effects, while undirected, mutual, and bidirected connections are converted into model covariances. SEMrun output contains different sets of parameter estimates. Beta coefficients (i.e., direct effects) are estimated from directed interactions and residual covariances (psi coefficients) from bidirected, undirected, or mutual interactions. If a group variable is given, exogenous group effects on nodes (gamma coefficients) or edges (delta coefficients) will be estimated. By default, maximum likelihood parameter estimates and P-values for parameter sets are computed by conventional z-test ( $= \text{estimate}/\text{SE}$ ), and fits it through the [lavaan](#) function, via Maximum Likelihood Estimation (`estimator = "ML"`, default estimator in [lavOptions](#)). In case of high dimensionality ( $n.\text{variables} \gg n.\text{subjects}$ ), the covariance matrix could not be semi-definite positive and thus parameter estimates could not be done. If this happens, covariance matrix regularization is enabled using the James-Stein-type shrinkage estimator implemented in the function [pcor.shrink](#) of `corpcor` R package. Argument `fit` determines how group influence is evaluated in the model, as absent (`fit = 0`), node perturbation (`fit = 1`), or edge perturbation (`fit = 2`). When `fit = 1`, the group is modeled as an exogenous variable, influencing all the other graph nodes. When `fit = 2`, SEMrun estimates the differences of the beta and/or psi coefficients (network edges) between groups. This is equivalent to fit a separate model for cases and controls, as opposed to one common model perturbed by the exogenous group effect. Once fitted, the two models are then compared to assess significant edge (i.e., direct effect) differences ( $d = \text{beta}_1 - \text{beta}_0$ ). P-values for parameter sets are computed by z-test ( $= d/\text{SE}$ ), through [lavaan](#). As an alternative to standard P-value calculation, SEMrun may use either RICF (randomization P-values) or GGM (de-sparsified P-values) methods. These algorithms are much faster than [lavaan](#) in case of large input graphs.

## Value

A list of 5 objects:

1. "fit", SEM fitted lavaan, ricf, or ggmncv object, depending on the MLE method specified by the `algo` argument;
2. "gest" or "dest", a data.frame of node-specific ("gest") or edge-specific ("dest") group effect estimates and P-values;
3. "model", SEM model as a string if `algo = "lavaan"`, and NULL otherwise;
4. "graph", the induced subgraph of the input network mapped on data variables. Graph edges (i.e., direct effects) with P-value  $< 0.05$  will be highlighted in red ( $\text{beta} > 0$ ) or blue ( $\text{beta} < 0$ ). If a group vector is given, nodes with significant group effect (P-value  $< 0.05$ ) will be red-shaded ( $\text{beta} > 0$ ) or lightblue-shaded ( $\text{beta} < 0$ );
5. "dataXY", input data subset mapping graph nodes, plus group at the first column (if no group is specified, this column will take NA values).

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

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**See Also**

See [fitAncestralGraph](#) and [fitConGraph](#) for RICF algorithm and constrained GGM algorithm details, respectively.

**Examples**

```
#### Model fitting (no group effect)

sem0 <- SEMrun(graph = sachs$graph, data = log(sachs$pkc))
summary(sem0$fit)
head(parameterEstimates(sem0$fit))

# Graphs
gplot(sem0$graph, main = "significant edge weights")
plot(sem0$graph, layout = layout.circle, main = "significant edge weights")

#### Model fitting (common model, group effect on nodes)

sem1 <- SEMrun(graph = sachs$graph, data = log(sachs$pkc),
               group = sachs$group)

# Fitting summaries
summary(sem1$fit)
```

```

print(sem1$gest)
head(parameterEstimates(sem1$fit))

# Graphs
gplot(sem1$graph, main = "Between group node differences")
plot(sem1$graph, layout = layout.circle, main = "Between group node differences")

#### Two-group model fitting (group effect on edges)

sem2 <- SEMrun(graph = sachs$graph, data = log(sachs$pkc),
               group = sachs$group,
               fit = 2)

# Summaries
summary(sem2$fit)
print(sem2$dest)
head(parameterEstimates(sem2$fit))

# Graphs
gplot(sem2$graph, main = "Between group edge differences")
plot(sem2$graph, layout = layout.circle, main = "Between group edge differences")

# Fitting and visualization of a large pathway:

g <- kegg.pathways[["MAPK signaling pathway"]]
G <- properties(g)[[1]]; summary(G)

library(huge)
als.npn <- huge.npn(alsData$exprs)

g1 <- SEMrun(G, als.npn, alsData$group, algo = "cggm")$graph
g2 <- SEMrun(g1, als.npn, alsData$group, fit = 2, algo = "cggm")$graph

# extract the subgraph with between group node and edge differences
g2 <- g2 - E(g2)[-which(E(g2)$color != "gray50")]
g <- properties(g2)[[1]]

# plot graph
E(g)$color <- E(g2)$color[E(g2) %in% E(g)]
gplot(g, l = "fdp", main = "node and edge group differences")

```



## Description

Four tree-based structure learning methods are implemented with graph and data-driven algorithms. A tree is an acyclic graph with  $p$  vertices and  $p-1$  edges. The graph method refers to the Steiner Tree (ST), a tree from an undirected graph that connects "seed" with additional nodes in the "most compact" way possible. The data-driven methods propose fast and scalable procedures based on Chu-Liu-Edmonds' algorithm (CLE) to recover a tree from a full graph. The first method, called Causal Additive Trees (CAT) uses pairwise mutual weights as input for CLE algorithm to recover a directed tree (an "arborescence"). The second one applies CLE algorithm for skeleton recovery and extends the skeleton to a tree (a "polytree") represented by a Completed Partially Directed Acyclic Graph (CPDAG). Finally, the Minimum Spanning Tree (MST) connecting an undirected graph (or on an undirected full graph) with minimal edge weights can be identified.

## Usage

```
SEMtree(
  graph,
  data,
  seed,
  type = "CAT",
  eweight = NULL,
  alpha = 0.05,
  verbose = FALSE,
  ...
)
```

## Arguments

graph	An igraph object.
data	A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes (variables).
seed	A vector of seed nodes.
type	Tree-based structure learning method. Four algorithms are available: <ul style="list-style-type: none"> <li>"ST". Steiner Tree (ST) identification via fast Kou's algorithm (Kou et al, 1981) connecting a set of seed nodes (called Terminal vertices) with connector nodes (called Steiner vertices) from input graph as defined in graph with minimal total distance on its edges. By default the edge weights are based on the pairwise correlation, <math>1 - \text{abs}(\text{cor}(j,k))</math>. If input graph has <math>E(\text{graph})\\$weight=1</math>, and <code>eweight = "custom"</code>, ST seeks a minimum subtree (i.e. subtree with minimal number of edges).</li> <li>"CAT" (default). Causal additive trees (CAT) algorithm as in Jakobsen et al. (2022). The argument <code>graph</code> is set to <code>NULL</code> (i.e., no input graph is needed). In the first step, a (univariate) generalized additive model (GAM) is employed to estimate the residual variances, <math>\text{var}(X(j) - [X(j) X(k)])</math> for all <math>j \neq k</math>, then use these to construct edge weights as inputs to the Chu-Liu-Edmonds' algorithm (Chow and Liu, 1968) to recover the arborescence. Argument <code>seed</code> must be specified to analyse a subset of nodes (variables) of interest.</li> </ul>

	<ul style="list-style-type: none"> <li>• "CPDAG". CLE algorithm for Skeleton Recovery and CPDAG estimation as in Lou et al. (2021). Together with "CAT" algorithm, "CPDAG" is data-driven and the argument graph is set to NULL. The key idea is to first recover the skeleton of the polytree by applying the CLE algorithm to the pairwise sample correlations of the data matrix. After the skeleton is recovered, the set of all v-structures can be correctly identified via a simple thresholding approach to pairwise sample correlations. CPDAG can be found applying iteratively only Rule 1 of Meek (1995). Argument seed must be specified to analyse a subset of nodes (variables) of interest.</li> <li>• "MST". Minimum Spanning Tree (MST) identification via Prim's algorithm (Prim, 1957). The latter finds the subset of edges that includes every vertex of the graph (as defined in graph) such that the sum of the weights of the edges can be minimized. The argument seed is set to NULL (i.e., no seed nodes are needed), or if argument seed is not NULL, argument graph is set to NULL to recover the MST of the full graph.</li> </ul>
eweight	Edge weight type for igraph object derived from <a href="#">weightGraph</a> or from user-defined distances. This option determines the weight-to-distance transform. If set to "NULL" (default), edge weights will be internally computed equal to $1 - \text{abs}(\text{cor})$ , i.e., $1 - \text{abs}(\text{pairwise Pearson's correlation})$ . If eweight = "kegg", repressing interactions (-1) will be set to 2 (maximum distance), neutral interactions (0) will be set to 1, and activating interactions (+1) will be set to 1 (minimum distance). If eweight = "zsign", all significant interactions will be set to 1 (minimum distance), while non-significant ones will be set to 2. If eweight = "pvalue", weights (p-values) will be transformed to the inverse of negative base-10 logarithm. If eweight = "custom", the algorithm will use the distance measure specified by the user as "weight" edge attribute.
alpha	Threshold for rejecting a pair of node being independent in "CPDAG" algorithm. The latter implements a natural v-structure identification procedure by thresholding the pairwise sample correlations over all adjacent pairs of edges with some appropriate threshold. By default, alpha = 0.05.
verbose	If TRUE, it shows the output tree (not recommended for large graphs).
...	Currently ignored.

## Details

If the input graph is a directed graph, ST and MST undirected trees are converted in directed trees using the [orientEdges](#) function.

## Value

An igraph object. If type = "ST", seed nodes are colored in "aquamarine" and connectors in "white". If type = "ST" and type = "MST", edges are colored in "green" if not present in the input, graph. If type = "CPDAG", bidirected edges are colored in "black" (if the algorithm is not able to establish the direction of the relationship between x and y).

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

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- Lou, X., Hu, Y., Li, X. (2022). Linear Polytree Structural Equation Models: Structural Learning and Inverse Correlation Estimation. *arXiv*: <<https://doi.org/10.48550/arxiv.2107.10955>>

## Examples

```
library(huge)
data <- huge.npn(alsData$exprs)
graph <- alsData$graph

# graph-based trees
seed <- V(graph)$name[sample(1:vcount(graph), 10)]
tree1<- SEMtree(graph, data, seed=seed, type="ST", verbose=TRUE)
tree2<- SEMtree(graph, data, seed=NULL, type="MST", verbose=TRUE)

# data-driven trees
V <- colnames(data)[colnames(data) %in% V(graph)$name]
tree3<- SEMtree(NULL, data, seed=V, type="CAT", verbose=TRUE)
tree4<- SEMtree(NULL, data, seed=V, type="CPDAG", alpha=0.05, verbose=TRUE)
```

---

Shipley.test

---

Missing edge testing implied by a graph with Shipley's basis-set

---

## Description

Compute all the P-values of the d-separation tests implied by the missing edges of a given acyclic graph (DAG). The conditioning set  $Z$  is represented, in a DAG, by the union of the parent sets of  $X$  and  $Y$  (Shipley, 2000). The results of every test, in a DAG, is then combined using the Fisher's statistic in an overall test of the fitted model  $C = -2 \cdot \sum(\log(P\text{-value}(k)))$ , where  $C$  is distributed as a chi-squared variate with  $df = 2k$ , as suggested by Shipley (2000).

**Usage**

```
Shipley.test(graph, data, MCX2 = FALSE, limit = 30000, verbose = TRUE, ...)
```

**Arguments**

<code>graph</code>	A directed graph as an <code>igraph</code> object.
<code>data</code>	A data matrix with subjects as rows and variables as columns.
<code>MCX2</code>	If TRUE, a Monte Carlo P-value of the combined C test is enabled.
<code>limit</code>	An integer value corresponding to the number of missing edges of the extracted acyclic graph. Beyond this limit, multicore computation is enabled to reduce the computational burden. By default, <code>limit = 30000</code> .
<code>verbose</code>	If TRUE, Shipley's test results will be showed to screen (default = TRUE).
<code>...</code>	Currently ignored.

**Value**

A list of three objects: (i) "dag": the DAG used to perform the Shipley test (ii) "dsep": the data.frame of all d-separation tests over missing edges in the DAG and (iii) "ctest": the overall Shipley's P-value.

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**References**

Shipley B (2000). A new inferential test for path models based on DAGs. *Structural Equation Modeling*, 7(2): 206-218. <[https://doi.org/10.1207/S15328007SEM0702\\_4](https://doi.org/10.1207/S15328007SEM0702_4)>

**Examples**

```
#\donttest{

library(huge)
als.npn <- huge.npn(alsData$exprs)

sem <- SEMrun(alsData$graph, als.npn)
C_test <- Shipley.test(sem$graph, als.npn, MCX2 = FALSE)
#MC_test <- Shipley.test(sem$graph, als.npn, MCX2 = TRUE)

#}
```

---

summary.GGM	<i>GGM model summary</i>
-------------	--------------------------

---

**Description**

Generate a summary for a constrained Gaussian Graphical Model (GGM) similar to lavaan-formatted summary

**Usage**

```
## S3 method for class 'GGM'  
summary(object, ...)
```

**Arguments**

object	A constrained GGM fitted model object.
...	Currently ignored.

**Value**

Shown the lavaan-formatted summary to console

**Author(s)**

Mario Grassi <mario.grassi@unipv.it>

**See Also**

[SEMrun](#).

**Examples**

```
sem0 <- SEMrun(sachs$graph, log(sachs$pkc), algo = "cggm")  
summary(sem0$fit)
```

---

summary.RICF	<i>RICF model summary</i>
--------------	---------------------------

---

**Description**

Generate a summary for a RICF solver similar to lavaan-formatted summary

**Usage**

```
## S3 method for class 'RICF'  
summary(object, ...)
```

Arguments

object            A RICF fitted model object.  
...               Currently ignored.

Value

Shown the lavaan-formatted summary to console

Author(s)

Mario Grassi <mario.grassi@unipv.it>

See Also

[SEMrun](#).

Examples

```
sem1 <- SEMrun(sachs$graph, log(sachs$pkc), sachs$group, algo = "ricf")
summary(sem1$fit)
```

---

weightGraph	<i>Graph weighting methods</i>
-------------	--------------------------------

---

Description

Add data-driven edge and node weights to the input graph.

Usage

```
weightGraph(graph, data, group = NULL, method = "r2z", limit = 10000, ...)
```

Arguments

graph            An igraph object.

data             A matrix or data.frame. Rows correspond to subjects, and columns to graph nodes.

group            Binary vector. This vector must be as long as the number of subjects. Each vector element must be 1 for cases and 0 for control subjects. By default, group = NULL. If group is not NULL, also node weighting is activated, and node weights correspond to the sign (-1 if  $z < -2$ , +1 if  $z > 2$ , 0 otherwise) and P-value of the z-test =  $b/SE(b)$  from simple linear regression  $y \sim x$  (i.e., `glm(node ~ group)`).

method           Edge weighting method. It can be one of the following:

1. "r2z", weight edges of a graph using Fisher's r-to-z transform (Fisher, 1915) to test the correlation coefficient of pairs of interacting nodes, if `group == NULL`. Otherwise, the correlation difference between group will be tested and edge weights correspond to the sign (-1 if  $z < -2$ , +1 if  $z > 2$ , 0 otherwise) and P-value of the group correlation difference.
2. "sem", edge weights are defined by a SEM model that implies testing the group effect simultaneously on the source node and the sink node. A new parameter `w` is defined as the weighted sum of the total effect of the group on source and sink nodes, adjusted by node degree centrality, and edge weights correspond to the sign (-1 if  $z < -2$ , +1 if  $z > 2$ , 0 otherwise) and P-value of the z-test =  $w/SE(w)$ . Not available if `group == NULL`.
3. "cov", edge weights are defined by a new parameter `w` combining the group effect on the source node (mean group difference, adjusted by source degree centrality), the sink node (mean group difference, adjusted by sink degree centrality), and the source-sink interaction (correlation difference). Edge weights correspond to the sign (-1 if  $z < -2$ , +1 if  $z > 2$ , 0 otherwise) and P-value of the z-test =  $w/SE(w)$  of the combined difference of the group over source node, sink node, and their connection. Not available if `group == NULL`.
4. "cfa", edge weights are defined by a CFA1 model that implies testing the group effect, `w` on a latent variable (LV) with observed indicators two interacting nodes, fixing loading coefficients and residual variances for model identification. Edge weights correspond to the sign (-1 if  $z < -2$ , +1 if  $z > 2$ , 0 otherwise) and P-value of the z-test =  $w/SE(w)$  of the group effect on the LV. Not available if `group == NULL`.

`limit` An integer value corresponding to the number of graph edges. Beyond this limit, multicore computation is enabled to reduce the computational burden. By default, `limit = 10000`.

`...` Currently ignored.

## Value

A weighted graph, as an `igraph` object.

## Author(s)

Mario Grassi <mario.grassi@unipv.it>

## References

Grassi M, Palluzzi F, Tarantino B (2022). SEMgraph: An R Package for Causal Network Analysis of High-Throughput Data with Structural Equation Models. *Bioinformatics*, 38 (20), 4829–4830 <<https://doi.org/10.1093/bioinformatics/btac567>>

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**Examples**

```
# Graph weighting
G <- weightGraph(graph = sachs$graph,
                 data = log(sachs$pkc),
                 group = sachs$group,
                 method = "r2z")

# New edge attributes
head(E(G)$pv); summary(E(G)$pv)
head(E(G)$zsign); table(E(G)$zsign)

# New nodes attributes
head(V(G)$pv); summary(V(G)$pv)
head(V(G)$zsign); table(V(G)$zsign)
```



# Index

activeModule, 3  
alsData, 5  
ancestors (ancestry), 6  
ancestry, 6  
  
boot, 48  
  
cluster\_leading\_eigen, 52  
clusterGraph, 7, 10, 14, 16, 18, 30, 32, 55  
clusterScore, 7, 8, 8, 14, 16, 18  
colorGraph, 10  
corr2graph, 12  
cplot, 8, 10, 14  
  
dagitty2graph, 15  
descendants (ancestry), 6  
  
extractClusters, 16  
  
factor.analysis, 19  
fitAncestralGraph, 63  
fitConGraph, 50, 63  
  
glmnet, 53, 54  
gplot, 20  
graph2dag, 22, 52, 54  
graph2dagitty, 23  
graph2lavaan, 24  
gsa.extract, 24  
  
igraph, 7, 14  
impliedConditionalIndependencies, 29  
  
kegg, 26  
kegg.pathways, 27  
  
lavaan, 60, 62  
lavaan2graph, 28  
lavOptions, 62  
localCI.test, 29  
localTests, 29  
  
mergeNodes, 9, 30  
model.learn, 16, 18, 32  
modelSearch, 35, 55  
  
orientEdges, 38, 66  
  
p.adjust, 11, 13, 36, 41, 49, 51, 56–58  
pairwiseMatrix, 39  
parameterEstimates, 40  
parents (ancestry), 6  
pathFinder, 40  
pcor.shrink, 62  
properties, 42  
protoclust, 30  
  
resizeGraph, 35, 43  
  
sachs, 45  
sem.mediation, 46  
SEMace, 40, 41, 48, 55  
SEMbap, 50  
SEMdag, 33, 34, 44, 53  
SEMdci, 55  
SEMgsa, 24–26, 57  
SEMPath, 40, 59  
SEMrun, 10, 11, 18, 55, 60, 69, 70  
SEMtree, 64  
ShipleY.test, 50, 51, 67  
siblings (ancestry), 6  
SILGGM, 50  
summary.GGM, 69  
summary.RICF, 69  
  
weightGraph, 3, 22, 33, 35, 66, 70