

# Package ‘BDgraph’

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**Title** Bayesian Structure Learning in Graphical Models using Birth-Death MCMC

**Version** 2.69

**Description** Statistical tools for Bayesian structure learning in undirected graphical models for continuous, discrete, and mixed data. The package is implemented the recent improvements in the Bayesian graphical models' literature, including Mohammadi and Wit (2015) <[doi:10.1214/14-BA889](https://doi.org/10.1214/14-BA889)>, Mohammadi et al. (2021) <[doi:10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)>, and Mohammadi and Wit (2019) <[doi:10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)>.

**URL** <https://www.uva.nl/profile/a.mohammadi>

**Imports** igraph, ggplot2, pROC

**Suggests** ssgraph, huge, tmvtnorm, skimr, knitr, rmarkdown

**VignetteBuilder** knitr

**License** GPL (>= 2)

**Repository** CRAN

**NeedsCompilation** yes

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**Description**

The R package **BDgraph** provides statistical tools for Bayesian structure learning in undirected graphical models for continuous, count, binary, and mixed data. The package is implemented the recent improvements in the Bayesian graphical models' literature, including Mohammadi and Wit (2015), Mohammadi et al. (2017), Dobra and Mohammadi (2018), and Letac et al. (2018). The computationally intensive tasks of the package are implemented in parallel using **OpenMP** in C++

and interfaced with R, to speed up the computations. Besides, the package contains several functions for simulation and visualization, as well as several multivariate datasets taken from the literature.

### How to cite this package

To cite **BDgraph** in publications use:

Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)

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### References

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**See Also**

[bdgraph](#), [bdgraph.mpl](#), [bdgraph.sim](#), [compare](#), [rgwish](#)

**Examples**

```
## Not run:
library( BDgraph )

set.seed( 10 )

# Generating multivariate normal data from a 'scale-free' graph
data.sim <- bdgraph.sim( n = 100, p = 10, graph = "scale-free", vis = TRUE )

# Running algorithm based on GGMS
bdgraph.obj <- bdgraph( data = data.sim, iter = 5000 )

summary( bdgraph.obj )

# To compare the result with true graph
compare( data.sim, bdgraph.obj, main = c( "Target", "BDgraph" ), vis = TRUE )

# Confusion Matrix
conf.mat( actual = data.sim, pred = bdgraph.obj )

conf.mat.plot( actual = data.sim, pred = bdgraph.obj )

# Running algorithm based on GGMS and marginal pseudo-likelihood
bdgraph.mpl.obj <- bdgraph.mpl( data = data.sim, iter = 5000 )

summary( bdgraph.mpl.obj )

# Confusion Matrix
conf.mat( actual = data.sim, pred = bdgraph.mpl.obj )

conf.mat.plot( actual = data.sim, pred = bdgraph.mpl.obj )

# To compare the results of both algorithms with true graph
compare( data.sim, list( bdgraph.obj, bdgraph.mpl.obj ),
        main = c( "Target", "BDgraph", "BDgraph_mpl" ), vis = TRUE )

## End(Not run)
```

---

adj2link

---

*Extract links from an adjacency matrix*


---

**Description**

Extract links from an adjacency matrix or an object of classes "sim" from function [bdgraph.sim](#) and "graph" from function [graph.sim](#).

**Usage**

```
adj2link( adj )
```

**Arguments**

adj adjacency matrix corresponding to a graph structure in which  $a_{ij} = 1$  if there is a link between nodes  $i$  and  $j$ , otherwise  $a_{ij} = 0$ . It can be an object with S3 class "sim" from function `bdgraph.sim`. It can be an object with S3 class "graph" from function `graph.sim`.

**Value**

matrix corresponding to the extracted links from graph structure.

**Author(s)**

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**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
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**See Also**

[link2adj](#), [graph.sim](#)

**Examples**

```
# Generating a 'random' graph
adj <- graph.sim( p = 6, vis = TRUE )

adj2link( adj )
```

---

 bdgraph

*Search algorithm in graphical models*


---

### Description

As the main function of the **BDgraph** package, this function consists of several MCMC sampling algorithms for Bayesian model determination in undirected graphical models. To speed up the computations, the birth-death MCMC sampling algorithms are implemented in parallel using **OpenMP** in C++.

### Usage

```
bdgraph( data, n = NULL, method = "ggm", algorithm = "bdmcmc", iter = 5000,
         burnin = iter / 2, not.cont = NULL, g.prior = 0.5, df.prior = 3,
         g.start = "empty", jump = NULL, save = FALSE,
         cores = NULL, threshold = 1e-8, verbose = TRUE, nu = 1 )
```

### Arguments

data	there are two options: (1) an $(n \times p)$ matrix or a data.frame corresponding to the data, (2) an $(p \times p)$ covariance matrix as $S = X'X$ which $X$ is the data matrix ( $n$ is the sample size and $p$ is the number of variables). It also could be an object of class "sim", from function <a href="#">bdgraph.sim</a> . The input matrix is automatically identified by checking the symmetry.
n	number of observations. It is needed if the "data" is a covariance matrix.
method	character with two options "ggm" (default) and "gcg". Option "ggm" is for Gaussian graphical models based on Gaussianity assumption. Option "gcg" is for Gaussian copula graphical models for the data that not follow Gaussianity assumption (e.g. continuous non-Gaussian, count, or mixed dataset).
algorithm	character with two options "bdmcmc" (default) and "rjmc". Option "bdmcmc" is based on birth-death MCMC algorithm. Option "rjmc" is based on reversible jump MCMC algorithm. Option "bd-dmh" is based on birth-death MCMC algorithm using double Metropolis Hasting. Option "rj-dmh" is based on reversible jump MCMC algorithm using double Metropolis Hasting.
iter	number of iteration for the sampling algorithm.
burnin	number of burn-in iteration for the sampling algorithm.
not.cont	for the case method = "gcg", a vector with binary values in which 1 indicates not continuous variables.
g.prior	for determining the prior distribution of each edge in the graph. There are two options: a single value between 0 and 1 (e.g. 0.5 as a noninformative prior) or an $(p \times p)$ matrix with elements between 0 and 1.
df.prior	degree of freedom for G-Wishart distribution, $W_G(b, D)$ , which is a prior distribution of the precision matrix.

g.start	corresponds to a starting point of the graph. It could be an $(p \times p)$ matrix, "empty" (default), or "full". Option "empty" means the initial graph is an empty graph and "full" means a full graph. It also could be an object with S3 class "bdgraph" of R package <code>BDgraph</code> or the class " <code>ssgraph</code> " of R package <code>ssgraph::ssgraph()</code> ; this option can be used to run the sampling algorithm from the last objects of previous run (see examples).
jump	it is only for the BDMCMC algorithm ( <code>algorithm = "bdmcmc"</code> ). It is for simultaneously updating multiple links at the same time to update graph in the BDMCMC algorithm.
save	logical: if FALSE (default), the adjacency matrices are NOT saved. If TRUE, the adjacency matrices after burn-in are saved.
cores	number of cores to use for parallel execution. The case <code>cores = "all"</code> means all CPU cores to use for parallel execution.
threshold	threshold value for the convergence of sampling algorithm from G-Wishart for the precision matrix.
verbose	logical: if TRUE (default), report/print the MCMC running time.
nu	prior parameter for option <code>method = "tgm"</code> .

### Value

An object with S3 class "bdgraph" is returned:

p_links	upper triangular matrix which corresponds the estimated posterior probabilities of all possible links.
K_hat	posterior estimation of the precision matrix.

For the case `"save = TRUE"` is returned:

sample_graphs	vector of strings which includes the adjacency matrices of visited graphs after burn-in.
graph_weights	vector which includes the waiting times of visited graphs after burn-in.
all_graphs	vector which includes the identity of the adjacency matrices for all iterations after burn-in. It is needed for monitoring the convergence of the BD-MCMC algorithm.
all_weights	vector which includes the waiting times for all iterations after burn-in. It is needed for monitoring the convergence of the BD-MCMC algorithm.

### Author(s)

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### References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
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### See Also

[bdgraph.mpl](#), [bdgraph.dw](#), [bdgraph.sim](#), [summary.bdgraph](#), [compare](#)

### Examples

```
## Not run:
set.seed( 10 )

# - - Example 1

# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 100, p = 10, size = 15, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, iter = 1000, save = TRUE )

summary( bdgraph.obj )

# Confusion Matrix
conf.mat( actual = data.sim, pred = bdgraph.obj )

conf.mat.plot( actual = data.sim, pred = bdgraph.obj )

# To compare our result with true graph
compare( data.sim, bdgraph.obj, main = c( "Target", "BDgraph" ), vis = T )

# Running algorithm with starting points from previous run
bdgraph.obj2 <- bdgraph( data = data.sim, g.start = bdgraph.obj )

compare( data.sim, list( bdgraph.obj, bdgraph.obj2 ),
        main = c( "Target", "Frist run", "Second run" ) )

# - - Example 2

# Generating mixed data from a 'scale-free' graph
data.sim <- bdgraph.sim( n = 200, p = 7, type = "mixed", graph = "scale-free", vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, method = "gcgm" )

summary( bdgraph.obj )
```



```

compare( data.sim, bdgraph.obj, vis = T )

conf.mat( actual = data.sim, pred = bdgraph.obj )

conf.mat.plot( actual = data.sim, pred = bdgraph.obj )

## End(Not run)

```

---

bdgraph.dw	<i>Search algorithm for Gaussian copula graphical models for count data</i>
------------	---

---

## Description

This function consists of several sampling algorithms for Bayesian structure learning in undirected graphical models for count data. It is based on Gaussian copula graphical models with discrete Weibull distributed marginals. To speed up the computations, the birth-death MCMC sampling algorithms are implemented in parallel using **OpenMP** in C++.

## Usage

```

bdgraph.dw( data, x = NULL, formula = y ~ .,
            n = NULL, algorithm = "bdmcmc", iter = 5000,
            burnin = iter / 2, g.prior = 0.5, df.prior = 3,
            ZI = FALSE, iter_bdw = 5000,
            g.start = "empty", jump = NULL, save = FALSE,
            q = NULL, beta = NULL, pii = NULL,
            cores = NULL, threshold = 1e-8, verbose = TRUE )

```

## Arguments

data	$(n \times p)$ matrix or a data.frame corresponding to the data on the $p$ nodes of the graph. It can also be an object of class "sim", from the function <a href="#">bdgraph.sim</a> .
x	$(n \times k)$ matrix or a data.frame corresponding to the predictors.
formula	object of class <a href="#">formula</a> as a symbolic description of the model for linking each node to the predictors. For the case of data.frame, it is taken as the model frame (see <a href="#">model.frame</a> ).
n	number of observations. It is needed if the "data" is a covariance matrix.
algorithm	character with two options "bdmcmc" (default) and "rjmc". Option "bdmcmc" is based on a birth-death MCMC algorithm. Option "rjmc" is based on a reversible jump MCMC algorithm.
iter	number of iterations for the sampling algorithm for graph learning.
burnin	number of burn-in iterations for the sampling algorithm for graph learning.
g.prior	for determining the prior distribution of each edge in the graph. There are two options: a single value between 0 and 1 (e.g. 0.5 as a noninformative prior) or a $(p \times p)$ matrix with elements between 0 and 1.

<code>df.prior</code>	degree of freedom for G-Wishart distribution, $W_G(b, D)$ , which is a prior distribution for the precision matrix.
<code>ZI</code>	logical. If FALSE (default), the conditional distribution of each response variable is assumed to be Discrete Weibull given the predictors $x$ . If TRUE, a zero-inflated model will be applied to each response. <code>ZI</code> can be passed also as a vector, in order to specify which of the ( $p$ variables) should be fitted with zero-inflation (TRUE) or not (FALSE).
<code>iter_bdw</code>	number of iterations for the sampling algorithm to estimate the regression parameters for the Discrete Weibull distribution. It is passed to the <code>bdw.reg</code> function.
<code>g.start</code>	corresponds to a starting point of the graph. It could be an $(p \times p)$ matrix, "empty" (default), or "full". Option "empty" means that the initial graph is an empty graph and "full" means a full graph. It also could be an object with S3 class "bdgraph" of R package <code>BDgraph</code> or the class "ssgraph" of R package <code>ssgraph::ssgraph()</code> ; this option can be used to run the sampling algorithm from the last objects of the previous run (see examples).
<code>jump</code>	it is only for the BDMCMC algorithm ( <code>algorithm = "bdmcmc"</code> ). It is for simultaneously updating multiple links at the same time while updating the graph in the BDMCMC algorithm.
<code>save</code>	logical: if FALSE (default), the adjacency matrices are NOT saved. If TRUE, the adjacency matrices after burn-in are saved.
<code>q, beta</code>	parameters of the discrete Weibull distribution used for the marginals. They should be given either as a $(n \times p)$ matrix (if covariates are present) or as a vector (if covariates are not present). If NULL (default), these parameters are estimated by the <code>bdw.reg</code> function.
<code>pii</code>	vector of zero-inflation parameters of the zero-inflated discrete Weibull distributions used for the marginals. If NULL (default), this parameter is estimated by the <code>bdw.reg</code> function when <code>ZI = TRUE</code> .
<code>cores</code>	number of cores to use for parallel execution. The case <code>cores = "all"</code> means all CPU cores to use for parallel execution.
<code>threshold</code>	threshold value for the convergence of the sampling algorithm from G-Wishart for the precision matrix.
<code>verbose</code>	logical: if TRUE (default), report/print the MCMC running time.

### Value

An object with S3 class "bdgraph" is returned, containing:

<code>p_links</code>	upper triangular matrix corresponding to the estimated posterior probabilities of all possible links.
<code>K_hat</code>	posterior estimation of the precision matrix.
<code>sample_marginals</code>	posterior samples of the regression coefficients of the marginal distributions.

For the case `"save = TRUE"`, the code returns:

sample_graphs	vector of strings which includes the adjacency matrices of the graphs visited after burn-in.
graph_weights	vector which includes the waiting times of the graphs visited after burn-in.
all_graphs	vector which includes the identity of the adjacency matrices for all iterations after burn-in. It is needed for monitoring the convergence of the BDMCMC algorithm.
all_weights	vector which includes the waiting times for all iterations after burn-in. It is needed for monitoring the convergence of the BDMCMC algorithm.

### Author(s)

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### References

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- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377

### See Also

[bdgraph](#), [bdgraph.mpl](#), [bdw.reg](#), [bdgraph.sim](#), [summary.bdgraph](#), [compare](#)

### Examples

```
## Not run:
# - - Example 1

# Generating multivariate Discrete Weibull data based on 'random' graph
data.sim <- bdgraph.sim( n = 100, p = 10, type = "dw", vis = TRUE )

bdgraph.obj <- bdgraph.dw( data = data.sim, iter = 5000 )
```

```

summary( bdgraph.obj )

# To compare the result with true graph
compare( data.sim, bdgraph.obj, main = c( "Target", "BDgraph" ), vis = TRUE )

# - - Example 2

# Generating multivariate Discrete Weibull data based on a 'scale-free' graph
data.sim <- bdgraph.sim( n = 100, p = 10, type = "dw", graph = "scale-free", vis = TRUE )

bdgraph.obj <- bdgraph.dw( data = data.sim, iter = 10000 )

summary( bdgraph.obj )

compare( data.sim, bdgraph.obj, main = c( "Target", "BDgraph" ), vis = TRUE )

## End(Not run)

```

---

bdgraph.mpl                      *Search algorithm in graphical models using marginal pseudo-likelihood*

---

## Description

This function consists of several sampling algorithms for Bayesian model determination in undirected graphical models based on marginal pseudo-likelihood. To speed up the computations, the birth-death MCMC sampling algorithms are implemented in parallel using **OpenMP** in C++.

## Usage

```

bdgraph.mpl( data, n = NULL, method = "ggm", transfer = TRUE,
             algorithm = "bdmcmc", iter = 5000, burnin = iter / 2,
             g.prior = 0.5, g.start = "empty",
             jump = NULL, alpha = 0.5, save = FALSE,
             cores = NULL, operator = "or", verbose = TRUE )

```

## Arguments

data	there are two options: (1) an $(n \times p)$ matrix or a <code>data.frame</code> corresponding to the data, (2) an $(p \times p)$ covariance matrix as $S = X'X$ which $X$ is the data matrix ( $n$ is the sample size and $p$ is the number of variables). It also could be an object of class "sim", from function <code>bdgraph.sim</code> . The input matrix is automatically identified by checking the symmetry.
n	number of observations. It is needed if the "data" is a covariance matrix.
method	character with two options "ggm" (default), "dgm" and "dgm-binary". Option "ggm" is for Gaussian graphical models based on Gaussianity assumption. Option "dgm" is for discrete graphical models for the count data. Option "dgm-binary" is for discrete graphical models for the data that are binary.

transfer	for only 'count' data which method = "dgm" or method = "dgm-binary".
algorithm	character with two options "bdmcmc" (default) and "rjmcmc". Option "bdmcmc" is based on birth-death MCMC algorithm. Option "rjmcmc" is based on reversible jump MCMC algorithm. Option "hc" is based on hill-climbing algorithm; this algorithm is only for count data which method = "dgm" or method = "dgm-binary".
iter	number of iteration for the sampling algorithm.
burnin	number of burn-in iteration for the sampling algorithm.
g.prior	for determining the prior distribution of each edge in the graph. There are two options: a single value between 0 and 1 (e.g. 0.5 as a noninformative prior) or an $(p \times p)$ matrix with elements between 0 and 1.
g.start	corresponds to a starting point of the graph. It could be an $(p \times p)$ matrix, "empty" (default), or "full". Option "empty" means the initial graph is an empty graph and "full" means a full graph. It also could be an object with S3 class "bdgraph" of R package <a href="#">BDgraph</a> or the class "ssgraph" of R package <a href="#">ssgraph::ssgraph()</a> ; this option can be used to run the sampling algorithm from the last objects of previous run (see examples).
jump	it is only for the BDMCMC algorithm (algorithm = "bdmcmc"). It is for simultaneously updating multiple links at the same time to update graph in the BDMCMC algorithm.
alpha	value of the hyper parameter of Dirichlet, which is a prior distribution.
save	logical: if FALSE (default), the adjacency matrices are NOT saved. If TRUE, the adjacency matrices after burn-in are saved.
cores	number of cores to use for parallel execution. The case cores = "all" means all CPU cores to use for parallel execution.
operator	character with two options "or" (default) and "and". It is for hill-climbing algorithm.
verbose	logical: if TRUE (default), report/print the MCMC running time.

### Value

An object with S3 class "bdgraph" is returned:

`p_links` upper triangular matrix which corresponds the estimated posterior probabilities of all possible links.

For the case "save = TRUE" is returned:

`sample_graphs` vector of strings which includes the adjacency matrices of visited graphs after burn-in.

`graph_weights` vector which includes the waiting times of visited graphs after burn-in.

`all_graphs` vector which includes the identity of the adjacency matrices for all iterations after burn-in. It is needed for monitoring the convergence of the BD-MCMC algorithm.

`all_weights` vector which includes the waiting times for all iterations after burn-in. It is needed for monitoring the convergence of the BD-MCMC algorithm.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>, Adrian Dobra, and Johan Pensar

**References**

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**See Also**

[bdgraph](#), [bdgraph.dw](#), [bdgraph.sim](#), [summary.bdgraph](#), [compare](#)

**Examples**

```
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 70, p = 5, size = 7, vis = TRUE )

bdgraph.obj <- bdgraph.mpl( data = data.sim, iter = 500 )

summary( bdgraph.obj )

# To compare the result with true graph
compare( data.sim, bdgraph.obj, main = c( "Target", "BDgraph" ) )
```

---

bdgraph.npn

*Nonparametric transfer*

---

**Description**

Transfers non-Gaussian data to Gaussian.

**Usage**

```
bdgraph.npn( data, npn = "shrinkage", npn.thresh = NULL )
```

**Arguments**

data	$(n \times p)$ matrix or a data.frame corresponding to the data ( $n$ is the sample size and $p$ is the number of variables).
npn	character with three options "shrinkage" (default), "truncation", and "skeptical". Option "shrinkage" is for the shrunken transformation, option "truncation" is for the truncated transformation and option "skeptical" is for the non-paranormal skeptical transformation. For more details see references.
npn.thresh	truncation threshold; it is only for the truncated transformation (npn= "truncation"). The default value is $1/(4n^{1/4}\sqrt{\pi \log(n)})$ .

**Value**

$(n \times p)$  matrix of transferred data, if npn = "shrinkage" or "truncation", and a non-paranormal correlation  $(p \times p)$  matrix, if npn = "skeptical".

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**References**

Liu, H., et al (2012). High Dimensional Semiparametric Gaussian Copula Graphical Models, *Annals of Statistics*, 40(4):2293-2326

Zhao, T. and Liu, H. (2012). The **huge** Package for High-dimensional Undirected Graph Estimation in R, *Journal of Machine Learning Research*, 13:1059-1062

**See Also**

[bdgraph.sim](#), [bdgraph](#), [bdgraph.mpl](#)

**Examples**

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 6, p = 4, size = 4 )

data <- ( data.sim $ data - 3 ) ^ 4
data

# Transfer the data by truncation
bdgraph.npn( data, npn = "truncation" )

# Transfer the data by shrunken
bdgraph.npn( data, npn = "shrunken" )

# Transfer the data by skeptical
bdgraph.npn( data, npn = "skeptical" )

## End(Not run)
```

## Description

Simulating multivariate distributions with different types of underlying graph structures, including "random", "cluster", "smallworld", "scale-free", "lattice", "hub", "star", "circle", "AR(1)", and "AR(2)". Based on the underlying graph structure, the function generates different types of *multivariate* data, including "Gaussian", "non-Gaussian", "categorical", "pois" (Poisson), "nbinom" (negative binomial), "dweibull" (discrete Weibull), "binary", "t" (t-distribution), "alternative-t", or "mixed" data. This function can be used also for simulating only graphs by setting the option  $n=0$  (default).

## Usage

```
bdgraph.sim( p = 10, graph = "random", n = 0, type = "Gaussian", prob = 0.2,
             size = NULL, mean = 0, class = NULL, cut = 4, b = 3,
             D = diag( p ), K = NULL, sigma = NULL,
             q = exp(-1), beta = 1, vis = FALSE, rewire = 0.05,
             range.mu = c( 3, 5 ), range.dispersion = c( 0.01, 0.1 ), nu = 1 )
```

## Arguments

p	number of variables (nodes).
graph	graph structure with options "random", "cluster", "smallworld", "scale-free", "lattice", "hub", "star", "circle", "AR(1)", and "AR(2)". It could also be an adjacency matrix corresponding to a graph structure (an upper triangular matrix in which $g_{ij} = 1$ if there is a link between nodes $i$ and $j$ , otherwise $g_{ij} = 0$ ).
n	number of samples required. Note that for the case $n = 0$ , only the graph is generated.
type	type of data with options "Gaussian" (default), "non-Gaussian", "categorical", "pois", "nbinom", "dweibull", "binary", and "mixed". For the option "Gaussian", data are generated from a multivariate normal distribution. For the option "non-Gaussian", data are transferred from a multivariate normal distribution to a continuous multivariate non-Gaussian distribution via Exponential marginals. For the option "categorical", data are transferred from a multivariate normal distribution to multivariate 'categorical' data. For the option "pois", data are transferred from a multivariate normal distribution to a multivariate Poisson distribution. For the option "nbinom", data are transferred from a multivariate normal distribution to a multivariate Negative Binomial distribution. For the option "dweibull", data are transferred from a multivariate normal distribution to a multivariate discrete Weibull distribution with parameters $q$ and $\beta$ . For the option "binary", data are generated directly from the joint distribution, in this case $p$ must be less than 17. For the option "mixed", data are transferred from a multivariate normal distribution to a mixture of 'categorical', 'non-Gaussian', 'binary' and 'Gaussian', respectively.



prob	if graph = "random", it is the probability that a pair of nodes has a link.
size	number of links in the true graph (graph size).
mean	vector specifying the mean of the variables.
class	if graph = "cluster", it is the number of classes.
cut	if type = "categorical", it is the number of categories for simulating 'categorical' data.
b	degree of freedom for G-Wishart distribution, $W_G(b, D)$ .
D	positive definite ( $p \times p$ ) "scale" matrix for G-Wishart distribution, $W_G(b, D)$ . The default is an identity matrix.
K	if graph = "fixed", it is a positive-definite symmetric matrix, corresponding to the true precision matrix.
sigma	if graph = "fixed", it is a positive-definite symmetric matrix corresponding to the true covariance matrix.
q, beta	if type = "dweibull", they are the parameters of the discrete Weibull distribution with density

$$p(x, q, \beta) = q^{x^\beta} - q^{(x+1)^\beta}, \quad \forall x = \{0, 1, 2, \dots\}.$$

They can be given either as a vector of length p or as an ( $n \times p$ ) matrix, e.g. if covariates are available and a regression model is used.

vis	visualize the true graph structure.
rewire	rewiring probability for smallworld network. Must be between 0 and 1.
range.mu, range.dispersion	if type = "nbinom", vector with two elements specifying the range of parameters for the Negative Binomial distribution.
nu	if type = "t" or "alternative-t", it is the parameter of the t distribution with density.

### Value

An object with S3 class "sim" is returned:

data	generated data as an ( $n \times p$ ) matrix.
sigma	covariance matrix of the generated data.
K	precision matrix of the generated data.
G	adjacency matrix corresponding to the true graph structure.

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>, Pariya Behrouzi, Veronica Vinciotti, Ernst Wit, and Alexander Christensen

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:10.1214/14BA889
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171
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- Pensar, J. et al (2017) Marginal pseudo-likelihood learning of discrete Markov network structures, *Bayesian Analysis*, 12(4):1195-215, doi:10.1214/16BA1032

## See Also

[graph.sim](#), [bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( p = 10, n = 50, prob = 0.3, vis = TRUE )

print( data.sim )

# Generating multivariate normal data from a 'hub' graph
data.sim <- bdgraph.sim( p = 6, n = 3, graph = "hub", vis = FALSE )

round( data.sim $ data, 2 )

# Generating mixed data from a 'hub' graph
data.sim <- bdgraph.sim( p = 8, n = 10, graph = "hub", type = "mixed" )

round( data.sim $ data, 2 )

# Generating only a 'scale-free' graph (with no data)
graph.sim <- bdgraph.sim( p = 8, graph = "scale-free" )

plot( graph.sim )

graph.sim $ G

## End(Not run)
```

---

bdw.reg *Bayesian estimation of (zero-inflated) Discrete Weibull regression*

---

## Description

Bayesian estimation of the parameters for Discrete Weibull (DW) regression. The conditional distribution of the response given the predictors is assumed to be DW with parameters  $q$  and  $\beta$ , dependent on the predictors, and, with an additional parameter  $\pi$  under zero inflation.

## Usage

```
bdw.reg( data, formula = NA, iter = 5000, burnin = NULL,
         dist.q = dnorm, dist.beta = dnorm,
         par.q = c( 0, 1 ), par.beta = c( 0, 1 ), par.pi = c( 1, 1 ),
         initial.q = NULL, initial.beta = NULL, initial.pi = NULL,
         ZI = FALSE, scale.proposal = NULL, adapt = TRUE, print = TRUE )
```

## Arguments

data	data.frame or matrix corresponding to the data, containing the variables in the model.
formula	object of class <a href="#">formula</a> as a symbolic description of the model to be fitted. For the case of data.frame, it is taken as the model frame (see <a href="#">model.frame</a> ).
iter	number of iterations for the sampling algorithm.
burnin	number of burn-in iterations for the sampling algorithm.
dist.q	Prior density for the regression coefficients associated to the parameter $q$ . The default is a Normal distribution ( <a href="#">dnorm</a> ). Any density function which has two parameters and can support the <code>log = TRUE</code> flag can be used, e.g. <a href="#">dnorm</a> , <a href="#">dlnorm</a> , <a href="#">dunif</a> etc.
dist.beta	Prior density for the regression coefficients associated to the parameter $\beta$ . The default is a Normal distribution ( <a href="#">dnorm</a> ). Any density function which has two parameters and can support the <code>log = TRUE</code> flag can be used, e.g. <a href="#">dnorm</a> , <a href="#">dlnorm</a> , <a href="#">dunif</a> etc.
par.q	vector of length two corresponding to the parameters of <code>dist.q</code> .
par.beta	vector of length two corresponding to the parameters of <code>dist.beta</code> .
par.pi	vector of length two corresponding to the parameters of the <a href="#">beta</a> prior density on $\pi$ .
initial.q, initial.beta, initial.pi	vector of initial values for the regression coefficients and for $\pi$ (if <code>ZI = TRUE</code> ).
ZI	logical: if <code>FALSE</code> (default), the conditional distribution of the response given the predictors is assumed to be DW with parameters $q$ and $\beta$ . If <code>TRUE</code> , a zero-inflated DW distribution will be applied.
scale.proposal	scale of the proposal function. Setting to lower values results in an increase in the acceptance rate of the sampler.

adapt	logical: if TRUE (default), the proposals will be adapted. If FALSE, no adapting will be applied.
print	logical: if TRUE (default), tracing information is printed.

### Details

The regression model uses a logit link function on  $q$  and a log link function on  $\beta$ , the two parameters of a DW distribution, with probability mass function given by

$$DW(y) = q^{y^\beta} - q^{(y+1)^\beta}, y = 0, 1, 2, \dots$$

For the case of zero inflation (ZI = TRUE), a zero-inflated DW is considered:

$$f(y) = (1 - pi)I(y = 0) + piDW(y)$$

where  $0 \leq pi \leq 1$  and  $I(y = 0)$  is an indicator for the point mass at zero for the response  $y$ .

### Value

sample	MCMC samples
q.est	posterior estimates of $q$
beta.est	posterior estimates of $\beta$
pi.est	posterior estimates of $pi$
accept.rate	acceptance rate of the MCMC algorithm

### Author(s)

Veronica Vinciotti, Reza Mohammadi <a.mohammadi@uva.nl>, and Pariya Behrouzi

### References

- Vinciotti, V., Behrouzi, P., and Mohammadi, R. (2022) Bayesian structural learning of microbiota systems from count metagenomic data, *arXiv preprint*, doi:10.48550/arXiv.2203.10118
- Peluso, A., Vinciotti, V., and Yu, K. (2018) Discrete Weibull generalized additive model: an application to count fertility, *Journal of the Royal Statistical Society: Series C*, 68(3):565-583, doi:10.1111/rssc.12311
- Haselimashhadi, H., Vinciotti, V. and Yu, K. (2018) A novel Bayesian regression model for counts with an application to health data, *Journal of Applied Statistics*, 45(6):1085-1105, doi:10.1080/02664763.2017.1342782
- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03

### See Also

[bdgraph.dw](#), [bdgraph](#), [ddweibull](#), [bdgraph.sim](#)

**Examples**

```

## Not run:
# - - Example 1

q    = 0.6
beta = 1.1
n    = 500

y = BDgraph::rdweibull( n = n, q = q, beta = beta )

output = bdw.reg( data = y, y ~ ., iter = 5000 )

output $ q.est
output $ beta.est

traceplot( output $ sample[ , 1 ], acf = T, pacf = T )
traceplot( output $ sample[ , 2 ], acf = T, pacf = T )

# - - Example 2

q    = 0.6
beta = 1.1
pii  = 0.8
n    = 500

y_dw = BDgraph::rdweibull( n = n, q = q, beta = beta )
z = rbinom( n = n, size = 1, prob = pii )
y = z * y_dw

output = bdw.reg( data = y, iter = 5000, ZI = TRUE )

output $ q.est
output $ beta.est
output $ pi.est

traceplot( output $ sample[ , 1 ], acf = T, pacf = T )
traceplot( output $ sample[ , 2 ], acf = T, pacf = T )
traceplot( output $ sample[ , 3 ], acf = T, pacf = T )

# - - Example 3

theta.q    = c( 0.1, -0.1, 0.34 ) # true parameter
theta.beta = c( 0.1, -0.15, 0.5 ) # true parameter

n = 500

x1 = runif( n = n, min = 0, max = 1.5 )
x2 = runif( n = n, min = 0, max = 1.5 )

reg_q = theta.q[ 1 ] + x1 * theta.q[ 2 ] + x2 * theta.q[ 3 ]
q      = 1 / ( 1 + exp( - reg_q ) )

```

```

reg_beta = theta.beta[ 1 ] + x1 * theta.beta[ 2 ] + x2 * theta.beta[ 3 ]
beta      = exp( reg_beta )

y = BDgraph::rdweibull( n = n, q = q, beta = beta )

data = data.frame( x1, x2, y )

output = bdw.reg( data, y ~. , iter = 5000 )

# - - Example 4

theta.q   = c( 1, -1, 0.8 ) # true parameter
theta.beta = c( 1, -1, 0.3 ) # true parameter
pii = 0.8

n = 500

x1 = runif( n = n, min = 0, max = 1.5 )
x2 = runif( n = n, min = 0, max = 1.5 )

reg_q = theta.q[ 1 ] + x1 * theta.q[ 2 ] + x2 * theta.q[ 3 ]
q      = 1 / ( 1 + exp( - reg_q ) )

reg_beta = theta.beta[ 1 ] + x1 * theta.beta[ 2 ] + x2 * theta.beta[ 3 ]
beta      = exp( reg_beta )

y_dw = BDgraph::rdweibull( n = n, q = q, beta = beta )
z     = rbinom( n = n, size = 1, prob = pii )
y     = z * y_dw

data = data.frame( x1, x2, y )

output = bdw.reg( data, y ~. , iter = 5000 )

## End(Not run)

```

---

bf

*Bayes factor between two graphs*


---

### Description

Compute the Bayes factor between the structure of two graphs.

### Usage

```
bf( num, den, bdgraph.obj, log = TRUE )
```

### Arguments

num, den	adjacency matrix corresponding to the true graph structure in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "graph" from function <a href="#">graph.sim</a> . It can be an object with S3 class "sim" from function <a href="#">bdgraph.sim</a> .
bdgraph.obj	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> . It also can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
log	character value. If TRUE the Bayes factor is given as log(BF).

### Value

single numeric value, the Bayes factor of the two graph structures num and den.

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>

### References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, [doi:10.18637/jss.v089.i03](#)
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- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, [doi:10.1214/18AOAS1164](#)

### See Also

[bdgraph](#), [bdgraph.mpl](#), [compare](#), [bdgraph.sim](#)

### Examples

```
## Not run:
# Generating multivariate normal data from a 'circle' graph
data.sim <- bdgraph.sim( n = 50, p = 6, graph = "circle", vis = TRUE )

# Running sampling algorithm
bdgraph.obj <- bdgraph( data = data.sim )

graph_1 <- graph.sim( p = 6, vis = TRUE )
```

```
graph_2 <- graph.sim( p = 6, vis = TRUE )  
  
bf( num = graph_1, den = graph_2, bdgraph.obj = bdgraph.obj )  
  
## End(Not run)
```

---

churn

*Churn data set*

---

### Description

The data can be downloaded from IBM Sample Data Sets. Customer *churn* occurs when customers stop doing business with a company, also known as customer attrition. The data set contains 3333 rows (customers) and 20 columns (features). The "Churn" column is our target which indicate whether customer churned (left the company) or not.

### Usage

```
data( churn )
```

### Format

The churn dataset, as a data frame, contains 3333 rows (customers) and 20 columns (variables/features). The 20 variables are:

- State: Categorical, for the 50 states and the District of Columbia.
- Account.Length: count, how long account has been active.
- Area.Code: Categorical.
- Int.l.Plan: Categorical, yes or no, international plan.
- VMail.Plan: Categorical, yes or no, voice mail plan.
- VMail.Message: Count, number of voice mail messages.
- Day.Mins: Continuous, minutes customer used service during the day.
- Day.Calls: Count, total number of calls during the day.
- Day.Charge: Continuous, total charge during the day.
- Eve.Mins: Continuous, minutes customer used service during the evening.
- Eve.Calls: Count, total number of calls during the evening.
- Eve.Charge: Continuous, total charge during the evening.
- Night.Mins: Continuous, minutes customer used service during the night.
- Night.Calls: Count, total number of calls during the night.
- Night.Charge: Continuous, total charge during the night.
- Intl.Mins: Continuous, minutes customer used service to make international calls.
- Intl.Calls: Count, total number of international calls.
- Intl.Charge: Continuous, total international charge.
- CustServ.Calls: Count, number of calls to customer service.
- Churn: Categorical, True or False. Indicator of whether the customer has left the company (True or False).



**References**

Larose, D. T. and Larose, C. D. (2014). Discovering knowledge in data: an introduction to data mining. *John Wiley & Sons*.

**Examples**

```
data( churn )
summary( churn )
```

---

compare	<i>Graph structure comparison</i>
---------	-----------------------------------

---

**Description**

This function provides several measures to assess the performance of the graphical structure learning.

**Usage**

```
compare( actual, pred, main = NULL, vis = FALSE )
```

**Arguments**

actual	adjacency matrix corresponding to the true graph structure in which $a_{ij} = 1$ if there is a link between nodes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "sim" from function <a href="#">bdgraph.sim</a> . It can be an object with S3 class "graph" from function <a href="#">graph.sim</a> .
pred	adjacency matrix corresponding to an estimated graph. It can be an object with S3 class "bdgraph" from function <a href="#">bdgraph</a> . It can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> . It can be an object of S3 class "select", from the function <a href="#">huge.select</a> of R package <a href="#">huge</a> . It also can be a list of above objects for comparing two or more different approaches.
main	character vector giving the names for the result table.
vis	logical: if TRUE, visualize the true graph and estimated graph structures.

**Value**

True positive	number of correctly estimated links.
True negative	number of true non-existing links which is correctly estimated.
False positive	number of links which they are not in the true graph, but are incorrectly estimated.
False negative	number of links which they are in the true graph, but are not estimated.

F1-score	weighted average of the "positive predictive" and "true positive rate". The F1-score value reaches its best value at 1 and worst score at 0.
Specificity	Specificity value reaches its best value at 1 and worst score at 0.
Sensitivity	Sensitivity value reaches its best value at 1 and worst score at 0.
MCC	Matthews Correlation Coefficients (MCC) value reaches its best value at 1 and worst score at 0.

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>, Antonio Abbruzzo, and Ivan Vujacic

### References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
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### See Also

[bdgraph](#), [bdgraph.mpl](#), [bdgraph.sim](#), [plotroc](#)

### Examples

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

# Running sampling algorithm based on GGMS
sample.ggm <- bdgraph( data = data.sim, method = "ggm", iter = 10000 )

# Comparing the results
compare( data.sim, sample.ggm, main = c( "True", "GGM" ), vis = TRUE )

# Running sampling algorithm based on GCGMs
sample.gcg <- bdgraph( data = data.sim, method = "gcm", iter = 10000 )

# Comparing GGM and GCGM methods
compare( data.sim, list( sample.ggm, sample.gcg ),
        main = c( "True", "GGM", "GCGM" ), vis = TRUE )
```

```
## End(Not run)
```

---

conf.mat	<i>Confusion Matrix</i>
----------	-------------------------

---

## Description

Create a Confusion Matrix.

## Usage

```
conf.mat( pred, actual, cutoff = 0.5, proportion = FALSE,
          dnn = c( "Prediction", "Actual" ), ... )
```

## Arguments

pred	adjacency matrix corresponding to an estimated graph. It can be an object with S3 class "bdgraph" from function <a href="#">bdgraph</a> . It can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
actual	adjacency matrix corresponding to the actual graph structure in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "sim" from function <a href="#">bdgraph.sim</a> . It can be an object with S3 class "graph" from function <a href="#">graph.sim</a> . It can be a factor, numeric or character vector of responses (true class), typically encoded with 0 (controls) and 1 (cases). Only two classes can be used in a ROC curve.
cutoff	cutoff value for the case that pred is vector of probabilities. The default is 0.5.
proportion	logical: FALSE (default) for a confusion matrix with number of cases. TRUE, for a confusion matrix with the proportion of cases.
dnn	names to be given to the dimensions in the result (the dimnames names).
...	further arguments to be passed to table.

## Value

the results of table on pred and actual.

## Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>

## See Also

[conf.mat.plot](#), [compare](#), [roc](#), [bdgraph](#)

## Examples

```
## Not run:
set.seed( 100 )

# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

# Running sampling algorithm based on GGMS
sample.ggm <- bdgraph( data = data.sim, method = "ggm", iter = 10000 )

# Confusion Matrix for GGM method
conf.mat( pred = sample.ggm, actual = data.sim )

## End(Not run)
```

---

conf.mat.plot

*Plot Confusion Matrix*

---

## Description

Plot a Confusion Matrix.

## Usage

```
conf.mat.plot( pred, actual, cutoff = 0.5, conf.level = 0, margin = 1,
              color = c( "#ff83a8", "#83ff9b" ), ... )
```

## Arguments

pred	adjacency matrix corresponding to an estimated graph. It can be an object with S3 class "bdgraph" from function <code>bdgraph</code> . It can be an object of S3 class "ssgraph", from the function <code>ssgraph::ssgraph()</code> of R package <code>ssgraph::ssgraph()</code> .
actual	adjacency matrix corresponding to the actual graph structure in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "sim" from function <code>bdgraph.sim</code> . It can be an object with S3 class "graph" from function <code>graph.sim</code> . It can be a factor, numeric or character vector of responses (true class), typically encoded with 0 (controls) and 1 (cases). Only two classes can be used in a ROC curve.
cutoff	cutoff value for the case that pred is vector of probabilities. The default is 0.5.
conf.level	confidence level used for the confidence rings on the odds ratios. Must be a single nonnegative number less than 1; if set to 0 (the default), confidence rings are suppressed.
margin	numeric vector with the margins to equate. Must be one of 1 (the default), 2, or <code>c(1, 2)</code> , which corresponds to standardizing the row, column, or both margins in each 2 by 2 table. Only used if <code>std</code> equals "margins".

color            vector of length 2 specifying the colors to use for the smaller and larger diagonals of each 2 by 2 table.

...              options to be passed to `fourfoldplot`.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**See Also**

[conf.mat](#), [compare](#), [roc](#), [bdgraph](#)

**Examples**

```
## Not run:
set.seed( 100 )

# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

# Running sampling algorithm based on GGMS
sample.ggm <- bdgraph( data = data.sim, method = "ggm", iter = 10000 )

# Confusion Matrix for GGM method
conf.mat.plot( pred = sample.ggm, actual = data.sim )

## End(Not run)
```

---

covariance

*Estimated covariance matrix*

---

**Description**

Provides the estimated covariance matrix.

**Usage**

```
covariance( bdgraph.obj, round = 2 )
```

**Arguments**

`bdgraph.obj`    object of S3 class "bdgraph", from function [bdgraph](#). It also can be an object of S3 class "ssgraph", from the function [ssgraph::ssgraph\(\)](#) of R package [ssgraph::ssgraph\(\)](#).

`round`            value for rounding all probabilities to the specified number of decimal places.

**Value**

matrix which corresponds the estimated covariance matrix.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)

**See Also**

[bdgraph](#), [precision](#), [plinks](#)

**Examples**

```
## Not run:
# Generating multivariate normal data from a 'circle' graph
data.sim <- bdgraph.sim( n = 70, p = 6, graph = "circle", vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim )

covariance( bdgraph.obj ) # Estimated covariance matrix

data.sim $ sigma # True covariance matrix

## End(Not run)
```

**Description**

Density, distribution function, quantile function and random generation for the discrete Weibull distribution (type I) with parameters  $q$  and  $\beta$ .

**Usage**

```

ddweibull( x, q = exp( -1 ), beta = 1, zero = TRUE )
pdweibull( x, q = exp( -1 ), beta = 1, zero = TRUE )
qdweibull( p, q = exp( -1 ), beta = 1, zero = TRUE )
rdweibull( n, q = exp( -1 ), beta = 1, zero = TRUE )

```

**Arguments**

**x** vector of quantiles.  
**p** vector of probabilities.  
**q, beta** shape and scale parameters, the latter defaulting to 1.  
**zero** logical; if TRUE (default), the support contains 0; FALSE otherwise.  
**n** number of observations. If `length(n) > 1`, the length is taken to be the number required.

**Details**

The discrete Weibull distribution has density given by

$$f(x) = q^{x^\beta} - q^{(x+1)^\beta}, x = 0, 1, 2, \dots$$

For the case `zero = FALSE`:

$$f(x) = q^{(x-1)^\beta} - q^{x^\beta}, x = 1, 2, \dots$$

Cumulative distribution function

$$F(x) = 1 - q^{(x+1)^\beta}$$

For the case `zero = FALSE`, `x+1` should be replaced by `x`.

**Value**

`ddweibull` gives the density, `pdweibull` gives the distribution function, `qdweibull` gives the quantile function, and `rdweibull` generates random values.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>, Pariya Behrouzi, Veronica Vinciotti

**References**

Nakagawa, T. and Osaki, S. (1975). The Discrete Weibull Distribution. *IEEE Transactions on Reliability*, R-24, 300-301, doi:[10.1109/TR.1975.5214915](https://doi.org/10.1109/TR.1975.5214915)

**See Also**

[dweibull](#), [bdw.reg](#), [bdgraph.dw](#)

### Examples

```
n = 1000
q = 0.4
beta = 0.8

set.seed( 7 )

rdw = rdweibull( n = n, q = q, beta = beta )

plot( prop.table( table( rdw ) ), type = "h", col = "gray50" )

x = 0:max( rdw )

lines( x, ddweibull( x = x, q = q, beta = beta ), type = "o", col = "blue", lwd = 2 )

hist( pdweibull( x = rdw, q = q, beta = beta ) )

plot( ecdf( rdw ) )
lines( x, pdweibull( x, q = q, beta = beta ), col = "blue", lwd = 2, type = "s" )
```

---

geneExpression

*Human gene expression dataset*

---

### Description

The dataset contains human gene expression of 100 transcripts (with unique Illumina TargetID) measured on 60 unrelated individuals.

### Usage

```
data( geneExpression )
```

### Format

The format is a matrix with 60 rows (number of individuals) and 100 column (number of transcripts).

### References

- Bhadra, A. and Mallick, B. K. (2013). Joint High Dimensional Bayesian Variable and Covariance Selection with an Application to eQTL Analysis, *Biometrics*, 69(2):447-457, doi:[10.1111/biom.12021](https://doi.org/10.1111/biom.12021)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)



**Examples**

```
data( geneExpression )

dim( geneExpression )
head( geneExpression )
```

gnorm

*Normalizing constant for G-Wishart***Description**

Calculates log of the normalizing constant of G-Wishart distribution based on the Monte Carlo method, developed by Atay-Kayis and Massam (2005).

**Usage**

```
gnorm( adj, b = 3, D = diag( ncol( adj ) ), iter = 100 )
```

**Arguments**

adj	adjacency matrix corresponding to the graph structure. It is an upper triangular matrix in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ .
b	degree of freedom for G-Wishart distribution, $W_G(b, D)$ .
D	positive definite ( $p \times p$ ) "scale" matrix for G-Wishart distribution, $W_G(b, D)$ . The default is an identity matrix.
iter	number of iteration for the Monte Carlo approximation.

**Details**

Log of the normalizing constant approximation using Monte Carlo method for a G-Wishart distribution,  $K \sim W_G(b, D)$ , with density:

$$Pr(K) = \frac{1}{I(b, D)} |K|^{(b-2)/2} \exp \left\{ -\frac{1}{2} \text{trace}(K \times D) \right\}.$$

**Value**

Log of the normalizing constant of G-Wishart distribution.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

## References

Atay-Kayis, A. and Massam, H. (2005). A monte carlo method for computing the marginal likelihood in nondecomposable Gaussian graphical models, *Biometrika*, 92(2):317-335, doi:10.1093/biomet/92.2.317

Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377

Uhler, C., et al (2018) Exact formulas for the normalizing constants of Wishart distributions for graphical models, *The Annals of Statistics* 46(1):90-118, doi:10.1214/17AOS1543

Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:10.1214/14BA889

## See Also

[rgwish](#), [rwish](#)

## Examples

```
## Not run:
# adj: adjacency matrix of graph with 3 nodes and 2 links
adj <- matrix( c( 0, 0, 1,
                 0, 0, 1,
                 0, 0, 0 ), 3, 3, byrow = TRUE )

gnorm( adj, b = 3, D = diag( 3 ) )

## End(Not run)
```

---

graph.sim

*Graph simulation*

---

## Description

Simulating undirected graph structures, including "random", "cluster", "scale-free", "lattice", "hub", "star", and "circle".

## Usage

```
graph.sim( p = 10, graph = "random", prob = 0.2, size = NULL, class = NULL, vis = FALSE,
           rewire = 0.05 )
```

## Arguments

**p** number of variables (nodes).

graph	undirected graph with options "random", "cluster", "smallworld", "scale-free", "lattice", "hub", "star", and "circle". It also could be an adjacency matrix corresponding to a graph structure (an upper triangular matrix in which $g_{ij} = 1$ if there is a link between nodes $i$ and $j$ , otherwise $g_{ij} = 0$ ).
prob	if graph = "random", it is the probability that a pair of nodes has a link.
size	number of links in the true graph (graph size).
class	if graph = "cluster", it is the number of classes.
vis	visualize the true graph structure.
rewire	rewiring probability for smallworld network. Must be between 0 and 1.

**Value**

The adjacency matrix corresponding to the simulated graph structure, as an object with S3 class "graph".

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl> and Alexander Christensen

**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
- Pensar, J. et al (2017) Marginal pseudo-likelihood learning of discrete Markov network structures, *Bayesian Analysis*, 12(4):1195-215, doi:[10.1214/16BA1032](https://doi.org/10.1214/16BA1032)

**See Also**

[bdgraph.sim](#), [bdgraph](#), [bdgraph.mpl](#)

**Examples**

```
# Generating a 'hub' graph
adj <- graph.sim( p = 8, graph = "scale-free" )

plot( adj )

adj
```

---

`link2adj`*Extract links from an adjacency matrix*

---

### Description

Extract links from an adjacency matrix or an object of classes "sim" from function `bdgraph.sim` and "graph" from function `graph.sim`.

### Usage

```
link2adj( link, p = NULL )
```

### Arguments

<code>link</code>	$(2 \times p)$ matrix or a data.frame corresponding to the links from the graph structure.
<code>p</code>	number of nodes of the graph.

### Value

An adjacency matrix corresponding to a graph structure in which  $a_{ij} = 1$  if there is a link between nodes  $i$  and  $j$ , otherwise  $a_{ij} = 0$ .

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>

### References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
- Pensar, J. et al (2017) Marginal pseudo-likelihood learning of discrete Markov network structures, *Bayesian Analysis*, 12(4):1195-215, doi:[10.1214/16BA1032](https://doi.org/10.1214/16BA1032)

**See Also**

[adj2link](#), [graph.sim](#)

**Examples**

```
# Generating a 'random' graph
adj <- graph.sim( p = 6, vis = TRUE )

link <- adj2link( adj )

link2adj( link, p = 6 )
```

---

pgraph

*Posterior probabilities of the graphs*

---

**Description**

Provides the estimated posterior probabilities for the most likely graphs or a specific graph.

**Usage**

```
pgraph( bdgraph.obj, number.g = 4, adj = NULL )
```

**Arguments**

bdgraph.obj	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> .
number.g	number of graphs with the highest posterior probabilities to be shown. This option is ignored if 'adj' is specified.
adj	adjacency matrix corresponding to a graph structure. It is an upper triangular matrix in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It also can be an object of S3 class "sim", from function <a href="#">bdgraph.sim</a> .

**Value**

selected_g	adjacency matrices which corresponding to the graphs with the highest posterior probabilities.
prob_g	vector of the posterior probabilities of the graphs corresponding to 'selected_g'.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:10.1214/14BA889
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171

## See Also

[bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 6, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, save = TRUE )

# Estimated posterior probability of the true graph
pgraph( bdgraph.obj, adj = data.sim )

# Estimated posterior probability of first and second graphs with highest probabilities
pgraph( bdgraph.obj, number.g = 2 )

## End(Not run)
```

---

plinks

*Estimated posterior link probabilities*

---

## Description

Provides the estimated posterior link probabilities for all possible links in the graph.

## Usage

```
plinks( bdgraph.obj, round = 2, burnin = NULL )
```

**Arguments**

bdgraph.obj	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> . It also can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
round	value for rounding all probabilities to the specified number of decimal places.
burnin	number of burn-in iteration to scape.

**Value**

An upper triangular matrix which corresponds the estimated posterior probabilities for all possible links.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, [doi:10.18637/jss.v089.i03](#)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, [doi:10.1214/14BA889](#)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, [doi:10.1080/01621459.2021.1996377](#)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, [doi:10.1214/18AOAS1164](#)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, [doi:10.1111/rssc.12171](#)

**See Also**

[bdgraph](#), [bdgraph.mpl](#)

**Examples**

```
## Not run:
# Generating multivariate normal data from a 'circle' graph
data.sim <- bdgraph.sim( n = 70, p = 6, graph = "circle", vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, iter = 10000 )

plinks( bdgraph.obj, round = 2 )

## End(Not run)
```

---

plot.bdgraph

*Plot function for S3 class "bdgraph"*


---

### Description

Visualizes structure of the selected graphs which could be a graph with links for which their estimated posterior probabilities are greater than 0.5 or graph with the highest posterior probability.

### Usage

```
## S3 method for class 'bdgraph'
plot( x, cut = 0.5, number.g = NULL, main = NULL,
      layout = igraph::layout_with_fr, vertex.size = 2, vertex.color = "orange",
      vertex.frame.color = "orange", vertex.label = NULL, vertex.label.dist = 0.5,
      vertex.label.color = "blue", edge.color = "lightblue", ... )
```

### Arguments

x	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> .
cut	threshold for including the links in the selected graph based on the estimated posterior probabilities of the links; See the examples.
number.g	number of graphs with the highest probabilities. This option works for the case running function <a href="#">bdgraph()</a> with option <code>save = TRUE</code> ; See the examples.
main	Graphical parameter (see <a href="#">plot</a> ).
layout	vertex placement which is according to R package <a href="#">igraph</a> ; For different layouts, see <a href="#">layout</a> of R package <a href="#">igraph</a> .
vertex.size	vertex size which is according to R package <a href="#">igraph</a> .
vertex.color	vertex color which is according to R package <a href="#">igraph</a> .
vertex.frame.color	vertex frame color which is according to R package <a href="#">igraph</a> .
vertex.label	vertex label. The default vertex labels are the vertex ids.
vertex.label.dist	vertex label distance which is according to R package <a href="#">igraph</a> .
vertex.label.color	vertex label color which is according to R package <a href="#">igraph</a> .
edge.color	edge color which is according to R package <a href="#">igraph</a> .
...	additional plotting parameters. For the complete list, see <a href="#">igraph.plotting</a> of R package <a href="#">igraph</a> .

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit



## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:10.1214/14BA889
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171

## See Also

[bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:
set.seed( 100 )

# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 100, p = 15, graph = "random", prob = 0.2, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim )

plot( bdgraph.obj )

bdgraph.obj <- bdgraph( data = data.sim, save = TRUE )

plot( bdgraph.obj, cut = 0.5 )

plot( bdgraph.obj, number.g = 4 )

## End(Not run)
```

---

plot.graph

*Plot function for S3 class "graph"*

---

## Description

Visualizes structure of the graph.

**Usage**

```
## S3 method for class 'graph'
plot( x, cut = 0.5, mode = "undirected", diag = FALSE, main = NULL,
      layout = igraph::layout_with_fr, vertex.size = 2, vertex.color = "orange",
      vertex.frame.color = "orange", vertex.label = NULL, vertex.label.dist = 0.5,
      vertex.label.color = "blue", edge.color = "lightblue", ... )
```

**Arguments**

x	object of S3 class "graph", from function <a href="#">graph.sim</a> .
cut	for the case where input 'x' is the object of class "bdgraph" or "ssgraph". Threshold for including the links in the selected graph based on the estimated posterior probabilities of the links.
mode	type of graph which is according to R package <a href="#">igraph</a> .
diag	logical which is according to R package <a href="#">igraph</a> .
main	graphical parameter (see plot).
layout	vertex placement which is according to R package <a href="#">igraph</a> ; For different layouts, see <a href="#">layout</a> of R package <a href="#">igraph</a> .
vertex.size	vertex size which is according to R package <a href="#">igraph</a> .
vertex.color	vertex color which is according to R package <a href="#">igraph</a> .
vertex.frame.color	vertex frame color which is according to R package <a href="#">igraph</a> .
vertex.label	vertex label. The default vertex labels are the vertex ids.
vertex.label.dist	vertex label distance which is according to R package <a href="#">igraph</a> .
vertex.label.color	vertex label color which is according to R package <a href="#">igraph</a> .
edge.color	edge color which is according to R package <a href="#">igraph</a> .
...	additional plotting parameters. For the complete list, see <a href="#">igraph.plotting</a> of R package <a href="#">igraph</a> .

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](#)
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Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164

Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171

Mohammadi, A. and Dobra, A. (2017). The R Package **BDgraph** for Bayesian Structure Learning in Graphical Models, *ISBA Bulletin*, 24(4):11-16

### See Also

[graph.sim](#), [bdgraph.sim](#), [plot.igraph](#)

### Examples

```
# Generating a 'scale-free' graph
adj <- graph.sim( p = 20, graph = "scale-free" )

plot( adj )
```

---

plot.sim	<i>Plot function for S3 class "sim"</i>
----------	---

---

### Description

Visualizes structure of the simulated graph for an object of S3 class "sim", from function [bdgraph.sim](#).

### Usage

```
## S3 method for class 'sim'
plot( x, ... )
```

### Arguments

x	object of S3 class "sim", from function <a href="#">bdgraph.sim</a> .
...	additional plotting parameters. See <a href="#">plot.graph</a> and for the complete list <a href="#">igraph.plotting</a> of R package <a href="#">igraph</a> .

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
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- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
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## See Also

[graph.sim](#), [bdgraph.sim](#), [plot.graph](#), [plot.igraph](#)

## Examples

```
set.seed( 10 )

# Generating synthetic multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 10, p = 15 )

plot( data.sim )
```

---

plotcoda

*Convergence plot*

---

## Description

Visualizes the cumulative occupancy fractions of all possible links in the graph. It can be used for monitoring the convergence of the sampling algorithms, BDMCMC and RJMCMC.

## Usage

```
plotcoda( bdgraph.obj, thin = NULL, control = TRUE, main = NULL,
          verbose = TRUE, ... )
```

## Arguments

`bdgraph.obj` object of S3 class "bdgraph", from function [bdgraph](#). It also can be an object of S3 class "ssgraph", from the function [ssgraph::ssgraph\(\)](#) of R package [ssgraph::ssgraph\(\)](#).

thin	option for getting fast result for a cumulative plot according to part of the iteration.
control	logical: if TRUE (default) and the number of nodes is greater than 15, then 100 links randomly is selected for visualization.
main	graphical parameter (see plot).
verbose	logical: if TRUE (default), report/print the calculation progress.
...	system reserved (no specific usage).

### Details

Note that a spending time for this function depends on the number of nodes.

For fast result, you can choose bigger value for the 'thin' option.

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>

### References

Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)

Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)

Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)

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Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)

### See Also

[bdgraph](#), [bdgraph.mpl](#), [traceplot](#)

### Examples

```
## Not run:
# Generating multivariate normal data from a 'circle' graph
data.sim <- bdgraph.sim( n = 50, p = 6, graph = "circle", vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, iter = 10000, burnin = 0 , save = TRUE )

plotcoda( bdgraph.obj )

## End(Not run)
```

plotroc

*ROC plot***Description**

Draws the receiver operating characteristic (ROC) curve according to the true graph structure for object of S3 class "bdgraph", from function `bdgraph`.

**Usage**

```
plotroc( actual, pred, cut = 20, smooth = FALSE, label = TRUE,
         AUC = TRUE, main = "ROC Curve" )
```

**Arguments**

actual	adjacency matrix corresponding to the true graph structure in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "sim" from function <code>bdgraph.sim</code> . It can be an object with S3 class "graph" from function <code>graph.sim</code> .
pred	upper triangular matrix corresponding to the estimated posterior probabilities for all possible links. It can be an object with S3 class "bdgraph" from function <code>bdgraph</code> . It can be an object of S3 class "ssgraph", from the function <code>ssgraph::ssgraph()</code> of R package <code>ssgraph::ssgraph()</code> . It can be an object of S3 class "select", from the function <code>huge.select</code> of R package <code>huge</code> . Options <code>est2</code> , <code>est3</code> and <code>est4</code> are for comparing two or more different approaches.
cut	number of cut points.
smooth	logical: for smoothing the ROC curve.
label	logical: for adding legend to the ROC plot.
AUC	logical: to report AUC.
main	overall title for the plot.

**Author(s)**

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**References**

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
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Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164

Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171

### See Also

[roc](#), [pROC::plot.roc\(\)](#), [pROC::auc\(\)](#), [bdgraph](#), [bdgraph.mpl](#), [compare](#)

### Examples

```
## Not run:
# To generate multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 30, p = 6, size = 7, vis = TRUE )

# To Run sampling algorithm
bdgraph.obj <- bdgraph( data = data.sim, iter = 10000 )

# To compare the results
plotroc( data.sim, bdgraph.obj )

# To compare the results based on CGGMs approach
bdgraph.obj2 <- bdgraph( data = data.sim, method = "gcm", iter = 10000 )

# To Compare the resultss
plotroc( data.sim, list( bdgraph.obj, bdgraph.obj2 ), label = FALSE )

## End(Not run)
```

---

precision

*Estimated precision matrix*

---

### Description

Provides the estimated precision matrix.

### Usage

```
precision( bdgraph.obj, round = 2 )
```

### Arguments

bdgraph.obj	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> . It also can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
round	value for rounding all probabilities to the specified number of decimal places.

**Value**

matrix which corresponds the estimated precision matrix.

**Author(s)**

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**References**

Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)

Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)

Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)

Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)

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**See Also**

[bdgraph](#), [covariance](#), [plinks](#)

**Examples**

```
## Not run:
# Generating multivariate normal data from a 'circle' graph
data.sim <- bdgraph.sim( n = 70, p = 6, graph = "circle", vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim )

precision( bdgraph.obj ) # Estimated precision matrix

data.sim $ K # True precision matrix

## End(Not run)
```



---

print.bdgraph                      *Print function for S3 class "bdgraph"*

---

## Description

Prints the information about the selected graph which could be a graph with links for which their estimated posterior probabilities are greater than 0.5 or graph with the highest posterior probability. It provides adjacency matrix, size and posterior probability of the selected graph.

## Usage

```
## S3 method for class 'bdgraph'  
print( x, ... )
```

## Arguments

x                      object of S3 class "bdgraph", from function [bdgraph](#).  
...                    system reserved (no specific usage).

## Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, [doi:10.18637/jss.v089.i03](#)
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- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, [doi:10.1111/rssc.12171](#)

## See Also

[bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:  
# Generating multivariate normal data from a 'random' graph  
data.sim <- bdgraph.sim( n = 50, p = 6, vis = TRUE )  
  
bdgraph.obj <- bdgraph( data = data.sim )  
  
print( bdgraph.obj )  
  
## End(Not run)
```

---

print.sim

*Print function for S3 class "sim"*

---

## Description

Prints the information about the type of data, the sample size, the graph type, the number of nodes, number of links and sparsity of the true graph.

## Usage

```
## S3 method for class 'sim'  
print( x, ... )
```

## Arguments

x                    object of S3 class "sim", from function `bdgraph.sim`.  
...                   system reserved (no specific usage).

## Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:[10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)

**See Also**

[graph.sim](#), [bdgraph.sim](#)

**Examples**

```
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 20, p = 10, vis = TRUE )

print( data.sim )
```

---

reinis

*Risk factors of coronary heart disease*

---

**Description**

The dataset consist of 6 discrete variables as the potential risk factors of coronary heart disease. The data collected from 1841 men employed of a car factory in Czechoslovakia (Reinis et al. 1981).

**Usage**

```
data( reinis )
```

**Format**

The format is a matrix with 1841 rows (number of individuals) and 6 column (number of variables).

**References**

Edwards and Havranek (1985). A fast procedure for model search in multidimensional contingency tables, *Biometrika*, 72:339-351

Reinis et al (1981). Prognostic significance of the risk profile in the prevention of coronary heart disease, *Bratis. lek. Listy*, 76:137-150

Mohammadi, A. and Dobra, A. (2017). The R Package **BDgraph** for Bayesian Structure Learning in Graphical Models, *ISBA Bulletin*, 24(4):11-16

**Examples**

```
data( reinis )

summary( reinis )
```

rgwish

*Sampling from G-Wishart distribution***Description**

Generates random matrices, distributed according to the G-Wishart distribution with parameters  $b$  and  $D$ ,  $W_G(b, D)$  with respect to the graph structure  $G$ . Note this function works for both non-decomposable and decomposable graphs.

**Usage**

```
rgwish( n = 1, adj = NULL, b = 3, D = NULL, threshold = 1e-8 )
```

**Arguments**

n	number of samples required.
adj	adjacency matrix corresponding to the graph structure which can be non-decomposable or decomposable. It should be an upper triangular matrix in which $a_{ij} = 1$ if there is a link between nodes $i$ and $j$ , otherwise $a_{ij} = 0$ . adj could be an object of class "graph", from function <code>graph.sim</code> . It also could be an object of class "sim", from function <code>bdgraph.sim</code> . It also could be an object of class "bdgraph", from functions <code>bdgraph.mpl</code> or <code>bdgraph</code> .
b	degree of freedom for G-Wishart distribution, $W_G(b, D)$ .
D	positive definite ( $p \times p$ ) "scale" matrix for G-Wishart distribution, $W_G(b, D)$ . The default is an identity matrix.
threshold	threshold value for the convergence of sampling algorithm from G-Wishart.

**Details**

Sampling from G-Wishart distribution,  $K \sim W_G(b, D)$ , with density:

$$Pr(K) \propto |K|^{(b-2)/2} \exp \left\{ -\frac{1}{2} \text{trace}(K \times D) \right\},$$

which  $b > 2$  is the degree of freedom and  $D$  is a symmetric positive definite matrix.

**Value**

A numeric array, say  $A$ , of dimension  $(p \times p \times n)$ , where each  $A[, , i]$  is a positive definite matrix, a realization of the G-Wishart distribution,  $W_G(b, D)$ . Note, for the case  $n = 1$ , the output is a matrix.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

## References

- Lenkoski, A. (2013). A direct sampler for G-Wishart variates, *Stat*, 2:119-128, doi:[10.1002/sta4.23](https://doi.org/10.1002/sta4.23)
- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
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- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)

## See Also

[gnorm](#), [rwish](#)

## Examples

```
# Generating a 'circle' graph as a non-decomposable graph
adj <- graph.sim( p = 5, graph = "circle" )
adj    # adjacency of graph with 5 nodes

sample <- rgwish( n = 1, adj = adj, b = 3, D = diag( 5 ) )
round( sample, 2 )

sample <- rgwish( n = 5, adj = adj )
round( sample, 2 )
```

---

rmvnorm

*Generate data from the multivariate Normal distribution*

---

## Description

Random generation function from the multivariate Normal distribution with mean equal to *mean* and covariance matrix *sigma*.

## Usage

```
rmvnorm( n = 10, mean = rep( 0, length = ncol( sigma ) ),
         sigma = diag( length( mean ) ) )
```

## Arguments

n	Number of observations.
mean	Mean vector, default is <code>rep(0, length = ncol(sigma))</code> .
sigma	positive definite covariance matrix, default is <code>diag(length(mean))</code> .

**Value**

A numeric matrix with rows equal to  $n$  and columns equal to  $length(mean)$ .

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**See Also**

[bdgraph.sim](#), [rwish](#), [rgwish](#)

**Examples**

```
mean <- c( 5, 20 )
sigma <- matrix( c( 4, 2,
                  2, 5 ), 2, 2 ) # covariance matrix

sample <- rmvnorm( n = 500, mean = mean, sigma = sigma )
plot( sample )
```

---

 roc

*Build a ROC curve*


---

**Description**

This function builds a ROC curve specifically for graph structure learning and returns a “roc” object, a list of class “roc”. This object can be printed, plotted, or passed to the functions [pROC::roc\(\)](#), [pROC::ci\(\)](#), [pROC::smooth.roc\(\)](#) and [pROC::coords\(\)](#). Additionally, two roc objects can be compared with [pROC::roc.test\(\)](#). This function is based on the [roc](#) function of R package pROC.

**Usage**

```
roc( pred, actual, auc = TRUE, smooth = FALSE, plot = FALSE, quiet = TRUE, ... )
```

**Arguments**

pred	adjacency matrix corresponding to an estimated graph. It can be an object with S3 class "bdgraph" from function <a href="#">bdgraph</a> . It can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> . It can be a <a href="#">numeric</a> or <a href="#">ordered</a> vector of the same length than actual, containing the predicted value of each observation.
actual	adjacency matrix corresponding to the actual graph structure in which $a_{ij} = 1$ if there is a link between notes $i$ and $j$ , otherwise $a_{ij} = 0$ . It can be an object with S3 class "sim" from function <a href="#">bdgraph.sim</a> . It can be an object with S3 class "graph" from function <a href="#">graph.sim</a> . It can be a factor, numeric or character vector of responses (true class), typically encoded with 0 (controls) and 1 (cases). Only two classes can be used in a ROC curve.

smooth	if TRUE, the ROC curve is passed to <code>smooth</code> to be smoothed.
auc	compute the area under the curve (AUC)? If TRUE (default), additional arguments can be passed to <code>pROC::auc()</code> .
plot	plot the ROC curve? If TRUE, additional arguments can be passed to <code>pROC::plot.roc()</code> .
quiet	if TRUE, turn off <code>messages</code> when direction and levels are auto-detected.
...	further arguments to be passed to <code>pROC::roc()</code> .

### Value

If the data contained any NA value and `na.rm=FALSE`, NA is returned. Otherwise, if `smooth=FALSE`, a list of class “roc” with the following fields:

auc	if called with <code>auc=TRUE</code> , a numeric of class “auc” as defined in <code>pROC::auc()</code> .
ci	if called with <code>ci=TRUE</code> , a numeric of class “ci” as defined in <code>pROC::ci()</code> .
response	the response vector. Patients whose response is not <code>%in%</code> levels are discarded. If NA values were removed, a <code>na.action</code> attribute similar to <code>na.omit</code> stores the row numbers.
predictor	predictor vector converted to numeric as used to build the ROC curve. Patients whose response is not <code>%in%</code> levels are discarded. If NA values were removed, a <code>na.action</code> attribute similar to <code>na.omit</code> stores the row numbers.
original.predictor, original.response	response and predictor vectors as passed in argument.
levels	levels of the response as defined in argument.
controls	predictor values for the control observations.
cases	predictor values for the cases.
percent	if the sensitivities, specificities and AUC are reported in percent, as defined in argument.
direction	direction of the comparison, as defined in argument.
fun.sesp	function used to compute sensitivities and specificities. Will be re-used in bootstrap operations.
sensitivities	sensitivities defining the ROC curve.
specificities	specificities defining the ROC curve.
thresholds	thresholds at which the sensitivities and specificities were computed. See below for details.
call	how the function was called. See <code>match.call</code> for more details.

If `smooth=TRUE` a list of class “smooth.roc” as returned by `pROC::smooth()`, with or without additional elements `auc` and `ci` (according to the call).

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl>

## References

Tom Fawcett (2006) “An introduction to ROC analysis”. *Pattern Recognition Letters* **27**, 861–874, doi:10.1016/j.patrec.2005.10.010

Xavier Robin, Natacha Turck, Alexandre Hainard, *et al.* (2011) “pROC: an open-source package for R and S+ to analyze and compare ROC curves”. *BMC Bioinformatics*, **7**, 77, doi:10.1186/1471-21051277.

## See Also

[plotroc](#), [pROC::plot.roc\(\)](#), [pROC::auc\(\)](#), [pROC::print.roc\(\)](#), [bdgraph](#), [bdgraph.mpl](#), [compare](#)

## Examples

```
## Not run:
set.seed( 5 )

# Generating multivariate normal data from a 'scale-free' graph
data.sim = bdgraph.sim( n = 200, p = 15, graph = "scale-free", vis = TRUE )

# Running BDMCMC algorithm
sample.bdmcmc = bdgraph( data = data.sim, algorithm = "bdmcmc", iter = 10000 )

# ROC curve for BDMCMC algorithm
roc.bdmcmc = BDgraph::roc( pred = sample.bdmcmc, actual = data.sim, plot = TRUE )

# Running RJMCMC algorithm
sample.rjmc = bdgraph( data = data.sim, algorithm = "rjmc", iter = 10000 )

# ROC curve for RJMCMC algorithm
roc.rjmc = BDgraph::roc( pred = sample.rjmc, actual = data.sim, plot = TRUE )

# ROC curve for both BDMCMC and RJMCMC algorithms
pROC::ggroc( list( BDMCMC = roc.bdmcmc, RJMCMC = roc.rjmc ) )

## End(Not run)
```

---

 rwish

*Sampling from Wishart distribution*


---

## Description

Generates random matrices, distributed according to the Wishart distribution with parameters  $b$  and  $D$ ,  $W(b, D)$ .

## Usage

```
rwish( n = 1, p = 2, b = 3, D = diag( p ) )
```



**Arguments**

n	number of samples required.
p	number of variables (nodes).
b	degree of freedom for Wishart distribution, $W(b, D)$ .
D	positive definite $(p \times p)$ "scale" matrix for Wishart distribution, $W(b, D)$ . The default is an identity matrix.

**Details**

Sampling from Wishart distribution,  $K \sim W(b, D)$ , with density:

$$Pr(K) \propto |K|^{(b-2)/2} \exp \left\{ -\frac{1}{2} \text{trace}(K \times D) \right\},$$

which  $b > 2$  is the degree of freedom and  $D$  is a symmetric positive definite matrix.

**Value**

A numeric array, say  $A$ , of dimension  $(p \times p \times n)$ , where each  $A[, , i]$  is a positive definite matrix, a realization of the Wishart distribution  $W(b, D)$ . Note, for the case  $n = 1$ , the output is a matrix.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

**References**

- Lenkoski, A. (2013). A direct sampler for G-Wishart variates, *Stat*, 2:119-128, [doi:10.1002/sta4.23](https://doi.org/10.1002/sta4.23)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, [doi:10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, [doi:10.1080/01621459.2021.1996377](https://doi.org/10.1080/01621459.2021.1996377)
- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, [doi:10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)

**See Also**

[gnorm](#), [rgwish](#)

**Examples**

```
sample <- rwish( n = 3, p = 5, b = 3, D = diag( 5 ) )
round( sample, 2 )
```

---

select	<i>Graph selection</i>
--------	------------------------

---

### Description

Provides the selected graph which, based on input, could be a graph with links for which their estimated posterior probabilities are greater than 0.5 (default) or a graph with the highest posterior probability; see examples.

### Usage

```
select( bdgraph.obj, cut = NULL, vis = FALSE )
```

### Arguments

bdgraph.obj	matrix in which each element response to the weight of the links. It can be an object of S3 class "bdgraph", from function <a href="#">bdgraph</a> . It can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
cut	threshold for including the links in the selected graph based on the estimated posterior probabilities of the links; see the examples.
vis	visualize the selected graph structure.

### Value

An adjacency matrix corresponding to the selected graph.

### Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

### References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](#)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](#)
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- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](#)

**See Also**

[bdgraph](#), [bdgraph.mpl](#)

**Examples**

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim )

select( bdgraph.obj )

bdgraph.obj <- bdgraph( data = data.sim, save = TRUE )

select( bdgraph.obj )

select( bdgraph.obj, cut = 0.5, vis = TRUE )

## End(Not run)
```

---

sparsity

*Compute the sparsity of a graph*

---

**Description**

Compute the sparsity of a graph/network or an object of class "graph" from function [graph.sim](#) or an object of class "sim" from function [bdgraph.sim](#).

**Usage**

```
sparsity( adj )
```

**Arguments**

adj adjacency matrix corresponding to a graph structure in which  $a_{ij} = 1$  if there is a link between nodes  $i$  and  $j$ , otherwise  $a_{ij} = 0$ . It can be an object with S3 class "graph" from function [graph.sim](#). It can be an object with S3 class "sim" from function [bdgraph.sim](#).

**Value**

value corresponding to the graph sparsity which is the proportion of the non-links (non-zero elements) in adj.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl>

## References

Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03

## See Also

[graph.sim](#), [adj2link](#), [link2adj](#)

## Examples

```
# Generating a 'random' graph
adj <- graph.sim( p = 10, graph = "random", prob = 0.4, vis = TRUE )

sparsity( adj )
```

---

summary.bdgraph	<i>Summary function for S3 class "bdgraph"</i>
-----------------	--

---

## Description

Provides a summary of the results for function [bdgraph](#).

## Usage

```
## S3 method for class 'bdgraph'
summary( object, round = 2, vis = TRUE, ... )
```

## Arguments

object	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> .
round	value for rounding all probabilities to the specified number of decimal places.
vis	visualize the results.
...	additional plotting parameters for the case vis = TRUE. See <a href="#">plot.graph</a> .

## Value

selected_g	adjacency matrix corresponding to the selected graph which has the highest posterior probability.
p_links	upper triangular matrix corresponding to the posterior probabilities of all possible links.
K_hat	estimated precision matrix.

## Author(s)

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:10.18637/jss.v089.i03
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:10.1214/14BA889
- Mohammadi, R., Massam, H. and Letac, G. (2021). Accelerating Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Journal of the American Statistical Association*, doi:10.1080/01621459.2021.1996377
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:10.1111/rssc.12171

## See Also

[bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim )

summary( bdgraph.obj )

bdgraph.obj <- bdgraph( data = data.sim, save = TRUE )

summary( bdgraph.obj )

summary( bdgraph.obj, vis = FALSE )

## End(Not run)
```

---

surveyData

*Labor force survey data*

---

## Description

The survey dataset concerns 1002 males in the U.S labor force, described by Hoff (2007). The seven observed variables which have been measured on various scales are as follow: the income (income), degree (degree), the number of children (children), parents income (pincome), parents degree (pdegree), number of parents children (pchildren), and age (age).

**Usage**

```
data( surveyData )
```

**Format**

The format is a matrix with 1002 rows (number of individuals) and 7 column (number of variables).

**References**

Hoff, P. (2007). Extending the rank likelihood for semiparametric copula estimation, *The Annals of Applied Statistics*, 1(1), 265-283.

Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)

**Examples**

```
data( surveyData )
summary( surveyData )
```

---

traceplot	<i>Trace plot of graph size</i>
-----------	---------------------------------

---

**Description**

Trace plot for graph size for the objects of S3 class "bdgraph", from function [bdgraph](#). It is a tool for monitoring the convergence of the sampling algorithms, BDMCMC and RJMCMC.

**Usage**

```
traceplot ( bdgraph.obj, acf = FALSE, pacf = FALSE, main = NULL, ... )
```

**Arguments**

bdgraph.obj	object of S3 class "bdgraph", from function <a href="#">bdgraph</a> . It also can be an object of S3 class "ssgraph", from the function <a href="#">ssgraph::ssgraph()</a> of R package <a href="#">ssgraph::ssgraph()</a> .
acf	visualize the autocorrelation functions for graph size.
pacf	visualize the partial autocorrelations for graph size.
main	graphical parameter (see plot).
...	system reserved (no specific usage).

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl> and Ernst Wit

## References

- Mohammadi, R. and Wit, E. C. (2019). **BDgraph**: An R Package for Bayesian Structure Learning in Graphical Models, *Journal of Statistical Software*, 89(3):1-30, doi:[10.18637/jss.v089.i03](https://doi.org/10.18637/jss.v089.i03)
- Mohammadi, A. and Wit, E. C. (2015). Bayesian Structure Learning in Sparse Gaussian Graphical Models, *Bayesian Analysis*, 10(1):109-138, doi:[10.1214/14BA889](https://doi.org/10.1214/14BA889)
- Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:[10.1214/18AOAS1164](https://doi.org/10.1214/18AOAS1164)
- Mohammadi, A. et al (2017). Bayesian modelling of Dupuytren disease by using Gaussian copula graphical models, *Journal of the Royal Statistical Society: Series C*, 66(3):629-645, doi:[10.1111/rssc.12171](https://doi.org/10.1111/rssc.12171)
- Mohammadi, A. and Dobra, A. (2017). The R Package **BDgraph** for Bayesian Structure Learning in Graphical Models, *ISBA Bulletin*, 24(4):11-16

## See Also

[plotcoda](#), [bdgraph](#), [bdgraph.mpl](#)

## Examples

```
## Not run:
# Generating multivariate normal data from a 'random' graph
data.sim <- bdgraph.sim( n = 50, p = 6, size = 7, vis = TRUE )

bdgraph.obj <- bdgraph( data = data.sim, iter = 10000, burnin = 0, save = TRUE )

traceplot( bdgraph.obj )

traceplot( bdgraph.obj, acf = TRUE, pacf = TRUE )

## End(Not run)
```

---

transfer

*transfer for count data*

---

## Description

Transfers count data, by counting the duplicated rows.

## Usage

```
transfer( r_data )
```

## Arguments

`r_data` ( $n \times p$ ) matrix or a `data.frame` corresponding to the data ( $n$  is the sample size and  $p$  is the number of variables).

**Value**

$(n \times p + 1)$  matrix of transferred data, in which the last column is the frequency of duplicated rows.

**Author(s)**

Reza Mohammadi <a.mohammadi@uva.nl> and Adrian Dobra

**References**

Mohammadi, A. and Dobra, A. (2017). The R Package **BDgraph** for Bayesian Structure Learning in Graphical Models, *ISBA Bulletin*, 24(4):11-16

Dobra, A. and Mohammadi, R. (2018). Loglinear Model Selection and Human Mobility, *Annals of Applied Statistics*, 12(2):815-845, doi:10.1214/18AOAS1164

**See Also**

[bdgraph.mpl](#), [bdgraph.sim](#)

**Examples**

```
# Generating multivariate binary data from a 'random' graph
data.sim <- bdgraph.sim( n = 12, p = 4, size = 4, type = "binary" )
r_data <- data.sim $ data
r_data

# Transfer the data
transfer( r_data )
```



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