

Package ‘cglasso’

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Description

Conditional graphical lasso estimator is an extension of the graphical lasso proposed to estimate the conditional dependence structure of a set of p response variables given q predictors. This package provides suitable extensions developed to study datasets with censored and/or missing values. Standard conditional graphical lasso is available as a special case. Furthermore, the package provides an integrated set of core routines for visualization, analysis, and simulation of datasets with censored and/or missing values drawn from a Gaussian graphical model. Details about the implemented models can be found in Augugliaro et al. (2020b) <[doi:10.1007/s11222-020-09945-7](https://doi.org/10.1007/s11222-020-09945-7)>, Augugliaro et al. (2020a) <[doi:10.1093/biostatistics/kxy043](https://doi.org/10.1093/biostatistics/kxy043)>, Yin et al. (2001) <[doi:10.1214/11-AOAS494](https://doi.org/10.1214/11-AOAS494)> and Stadler et al. (2012) <[doi:10.1007/s11222-010-9219-7](https://doi.org/10.1007/s11222-010-9219-7)>.

Imports methods, MASS

License GPL (>= 2)

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Description

Conditional graphical lasso (cglasso) estimator (Yin *and other*, 2011) is an extension of the graphical lasso (Yuan *and other*, 2007) proposed to estimate the conditional dependence structure of a set of p response variables given q predictors. This package provides suitable extensions developed to study datasets with censored and/or missing values (Augugliaro *and other*, 2020a and Augugliaro *and other*, 2020b). Standard conditional graphical lasso is available as a special case. Furthermore, the cglasso package provides an integrated set of core routines for visualization, analysis, and simulation of datasets with censored and/or missing values drawn from a Gaussian graphical model.

Details

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References

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AIC.cglasso

*Akaike Information Criterion***Description**

'AIC' computes the 'Akaike Information Criterion'.

Usage

```
## S3 method for class 'cglasso'
AIC(object, k = 2, mle, ...)
```

Arguments

object	an R object inheriting class 'cglasso', that is, the output of the model-fitting functions cglasso and cggm .
k	the <i>penalty</i> parameter to be used; the default k = 2 is the classical AIC.
mle	logical. TRUE if the measure of goodness-of-fit should be computed using the maximum likelihood estimates. Default depends on the class of the argument object: mle = FALSE for objects of class cglasso and mle = TRUE for objects of class cggm.
...	further arguments passed to cggm .

Details

'AIC' computes the following measure of goodness-of-fit (Ibrahim *and other*, 2008):

$$-2 \text{Q-function} + k \text{ df},$$

where k is the *penalty* parameter and df represents the number of unique non-zero parameters in the fitted model.

The values of the Q-function function are computed using [QFun](#). By default, for an object of class cglasso these values are computed using the penalized estimates whereas maximum likelihood estimates are used if the object is of class cggm (see argument 'mle' in [QFun](#)).

The Akaike Information Criterion (AIC) is returned by letting $k = 2$ (default value of the function AIC) whereas the 'Bayesian Information Criterion' (BIC) is returned by letting $k = \log(n)$, where n is the sample size.

Function AIC can be passed to the functions [select.cglasso](#) and [summary.cglasso](#) to select and print the optimal fitted model, respectively.

The function [plot.GoF](#) can be used to graphically evaluate the behaviour of the fitted models in terms of goodness-of-fit.

Value

'AIC' return an R object of S3 class "GoF", i.e., a named list containing the following components:

value_gof	a matrix storing the values of the measure of goodness-of-fit used to evaluate the fitted models.
df	a matrix storing the number of the estimated non-zero parameters.
dfB	a matrix storing the number of estimated non-zero regression coefficients.
dfTht	a matrix storing the number of estimated non-zero partial correlation coefficients.
value	a matrix storing the values of the Q-function.
n	the sample size.
p	the number of response variables.
q	the number of columns of the design matrix X used to fit the model.
lambda	the λ -values used to fit the model.
nlambda	the number of λ -values used.
rho	the ρ -values used to fit the model.
nrho	the number of ρ -values used.
type	a description of the computed measure of goodness-of-fit.
model	a description of the fitted model passed through the argument object.

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References

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- Wit, E., Heuvel, E. V. D., & Romeijn, J. W. (2012) <doi:10.1111/j.14679574.2012.00530.x>. All models are wrong...?: an introduction to model uncertainty. *Statistica Neerlandica*, 66(3), 217-236.

See Also

[BIC.cglasso](#), [cglasso](#), [cggm](#), [QFun](#), [plot.GoF](#) and [summary.cglasso](#)

Examples

```
set.seed(123)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
```

```

q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
AIC(out)

out.mle <- cggm(out, lambda.id = 3L, rho.id = 3L)
AIC(out.mle)

```

BIC.cglasso

Bayesian Information Criterion

Description

‘BIC’ computes the Bayesian Information Criterion.

Usage

```

## S3 method for class 'cglasso'
BIC(object, g = 0, type, mle, ...)

```

Arguments

object	an R object inheriting class ‘cglasso’, that is, the output of the model-fitting functions cglasso and cggm .
g	a value belonging to the interval $[0, 1]$. Classical BIC is returned by letting $g = 0$ (default value), whereas extended BIC corresponds to the case $g = 0.5$.
type	character; if g is not zero then the measure proposed in Foygel <i>and other</i> (2010) is returned by setting <code>type = "FD"</code> , otherwise (<code>type = "CC"</code>) returns the measure proposed in Chen <i>and other</i> (2008, 2012). See section ‘Details’ for more details.
mle	logical. TRUE if the measure of goodness-of-fit should be computed using the maximum likelihood estimates. Default depends on the class of the argument object: <code>mle = FALSE</code> for objects of class <code>cglasso</code> and <code>mle = TRUE</code> for objects of class <code>cggm</code> .
...	further arguments passed to the model-fitting function cggm .

Details

‘BIC’ computes the Bayesian Information Criterion (BIC) for models fitted by [cglasso](#) or [cggm](#). As proposed in Ibrahim *and other* (2008), BIC computes the measure of goodness-of-fit by replacing the log-likelihood function with the Q-function, that is, the function maximized in the M-Step of the EM-algorithm. The values of the Q-function are computed using [QFun](#). By default, for an object

of class `cglasso` these values are computed using the penalized estimates whereas, if the object has class `cggm`, maximum likelihood estimates are used (see argument `'mle'` in `QFun`).

By default, BIC computes the standard BIC measure ($\gamma = 0$):

$$-2 \text{Q-function} + \log(n) \text{ df},$$

where n is the sample size and `df` represents the number of unique non-zero parameters in the fitted model.

If $\gamma \neq 0$, the default depends on the number of predictors (q).

If $q = 0$, BIC computes the measure of goodness-of-fit proposed in Foygel *and other* (2010) (type = "FD"):

$$\text{eBIC} = -2 \text{QFun} + (\log n + 4 \gamma \log p) \text{ df},$$

where γ is a value belonging to the interval $[0, 1]$ and indexing the measure of goodness-of-fit.

If $q \neq 0$, BIC computes the measure of goodness-of-fit proposed in Chen *and other* (2008, 2012) (type = "CC"):

$$\text{eBIC} = -2 \text{QFun} + (\log n + 2 \gamma \log q) \text{ df},$$

BIC can be passed to the functions `select.cglasso` and `summary.cglasso` to select and print the optimal fitted model, respectively.

The function `plot.GoF` can be used to graphically evaluate the behaviour of the fitted models in terms of goodness-of-fit.

Value

'BIC' returns an R object of S3 class "GoF", i.e. a named list containing the following components:

<code>value_gof</code>	a matrix storing the values of the measure of goodness-of-fit used to evaluate the fitted models.
<code>df</code>	a matrix storing the number of the estimated non-zero parameters.
<code>dfB</code>	a matrix storing the number of estimated non-zero regression coefficients.
<code>dfTht</code>	a matrix storing the number of estimated non-zero partial correlation coefficients.
<code>value</code>	a matrix storing the values of the Q-function.
<code>n</code>	the sample size.
<code>p</code>	the number of response variables.
<code>q</code>	the number of columns of the design matrix X used to fit the model.
<code>lambda</code>	the λ -values used to fit the model.
<code>nlambda</code>	the number of λ -values used.
<code>rho</code>	the ρ -values used to fit the model.
<code>nrho</code>	the number of ρ -values used.
<code>type</code>	a description of the computed measure of goodness-of-fit.
<code>model</code>	a description of the fitted model passed through the argument object.

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References

Foygel, R. and Drton, M. (2010). Extended Bayesian Information Criteria for Gaussian Graphical Models. In: Lafferty, J., Williams, C., Shawe-taylor, J., Zemel, R.s. and Culott, A. (editors), *Advances in Neural Information Processing Systems 23*. pp. 604–612.

Chen, J. and Chen, Z. (2008) <doi:10.1093/biomet/asn034>. Extended Bayesian information criteria for model selection with large model spaces. *Biometrika*, Vol. 95(2), pp. 759–771.

Chen, J. and Chen, Z. (2012) <doi:10.5705/ss.2010.216>. Extended BIC for small-n-large-p sparse GLM. *Statistica Sinica*, Vol. 22, pp. 555–574.

Wit, E., Heuvel, E. V. D., & Romeijn, J. W. (2012) <doi:10.1111/j.14679574.2012.00530.x>. All models are wrong...?: an introduction to model uncertainty. *Statistica Neerlandica*, 66(3), 217-236.

See Also

[cglasso](#), [cggm](#), [AIC.cglasso](#), [QFun](#), [plot.GoF](#) and [summary.cglasso](#)

Examples

```
set.seed(123)

# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
BIC(out) # standard BIC measure
BIC(out, mle = TRUE, g = 0.5, type = "FD") # eBIC proposed in Foygel and other (2010)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
BIC(out) # standard BIC measure
BIC(out, mle = TRUE, g = 0.5, type = "FD") # eBIC proposed in Foygel and other (2010)
BIC(out, mle = TRUE, g = 0.5, type = "CC") # eBIC proposed in Chen and other (2008, 2010)
```

cggm	<i>Post-Hoc Maximum Likelihood Refitting of a Conditional Graphical Lasso</i>
------	---

Description

'cggm' is used to perform post-hoc maximum likelihood refitting of a selected conditional graphical lasso model with censored and/or missing values.

Usage

```
cggm(object, GoF = AIC, lambda.id, rho.id, tp.min = 1.0E-6, ntp = 100L,
      maxit.em = 1.0E+4, thr.em = 1.0E-3, maxit.bcd = 1.0E+5, thr.bcd = 1.0E-4,
      trace = 0L, ...)
```

Arguments

object	an R object of S3 class 'cglasso', that is, the output of the function <code>cglasso</code> .
GoF	a valid goodness-of-fit function, such as <code>AIC.cglasso</code> or <code>BIC.cglasso</code> , or an R object of class 'GoF'.
lambda.id	an optional integer used to identify a λ -value stored in object. See section 'Details' for more details.
rho.id	an optional integer used to identify a ρ -value stored in object. See section 'Details' for more details.
tp.min	the smallest λ and ρ value. See section 'Details' for more details.
ntp	integer; number of λ - and ρ -values used to compute the coefficients path. See section 'Details' for more details.
maxit.em	maximum number of iterations of the EM algorithm. Default is 1.0E+4.
thr.em	threshold for the convergence of the EM algorithm. Default value is 1.0E-4.
maxit.bcd	maximum number of iterations of the glasso algorithm. Default is 1.0E+5.
thr.bcd	threshold for the convergence of the glasso algorithm. Default is 1.0E-4.
trace	integer for printing information out as iterations proceed: trace = 0 no information is printed out on screen; trace = 1 minimal information is printed; trace = 2 detailed information is printed on screen.
...	the <i>penalty</i> parameter passed to the goodness-of-fit function (argument 'GoF').

Details

The model-fitting function `cggm` is used to obtain the maximum likelihood estimates of a censored Gaussian graphical model whose structure was found by penalised `cglasso` inference. That is, given a fitted `cglasso` model, identified along the path via a goodness-of-fit function (passed by `GoF`) or via the optional arguments `lambda.id` and `rho.id`, `cggm` computes the maximum likelihood estimates of the parameters with zero constraints associated to the given structure.

Maximum likelihood estimates are computed using `cglasso` as workhorse function, that is, `cggm` fits a sequence of `cglasso` models, with length equal to `ntp`, reducing λ and ρ until they are equal to the value `'min(tp.min, lambda.min, rho.min)'`, where `lambda.min` and `rho.min` are the smallest λ - and ρ -values stored in object. Maximum likelihood estimates are obtained from the last fitted `cglasso` model.

The model-fitting function `cggm` returns an R object inheriting the S3 class `'cglasso'`, for which all printing and plotting functions, designed for a `cglasso` object, can be used (see section `'See Also'`). Function `ShowStructure` can be used to show the structure of the package.

NOTE: if the user tries a problem when the sample size is not large enough, then increase the number of fitted `cglasso` models (argument `ntp`).

Value

`cggm` returns an object of S3 class `"cggm"`, i.e., a list containing the following components:

<code>call</code>	the call that produced this object.
<code>Yipt</code>	an array storing the <code>'working response matrix'</code> .
<code>B</code>	the maximum likelihood estimate of the regression coefficient matrix.
<code>mu</code>	the fitted expected values.
<code>R</code>	the <code>'working residuals'</code> matrix.
<code>S</code>	the <code>'working empirical covariance matrix'</code> .
<code>Sgm</code>	the maximum likelihood estimate of the covariance matrix.
<code>Tht</code>	the maximum likelihood estimate of the precision matrix.
<code>dfB</code>	the number of estimated non-zero regression coefficients. Only for internal purpose.
<code>dfTht</code>	the number of estimated non-zero (off-diagonal) partial correlation coefficients. Only for internal purpose.
<code>InfoStructure</code>	a named list whose elements are used to store the information about the estimated networks. Only for internal purpose.
<code>nit</code>	the number of EM steps.
<code>Z</code>	the <code>'datacggm'</code> object used to compute the censored graphical lasso estimator.
<code>nlambda</code>	Only for internal purpose.
<code>lambda</code>	the λ -value of the selected <code>cglasso</code> model.
<code>nrho</code>	Only for internal purpose.
<code>rho</code>	the ρ -value of the selected <code>cglasso</code> model.
<code>maxit.em</code>	maximum number of iterations of the EM algorithm.
<code>thr.em</code>	threshold for the convergence of the EM algorithm.
<code>maxit.bcd</code>	maximum number of iterations of the glasso algorithm.
<code>thr.bcd</code>	threshold for the convergence of the glasso algorithm.
<code>conv</code>	a description of the error that has occurred.
<code>subrout</code>	the name of the Fortran subroutine where the error has occurred (for internal debug only).

trace the integer used for printing information on screen.
nobs the sample size
nresp the number of response variables used to fit the model.
npred the number of predictors used to fit the model.

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See Also

[cglasso](#), [coef.cglasso](#), [fitted.cglasso](#), [residuals.cglasso](#), [predict.cggm](#), [impute](#), [AIC.cglasso](#), [BIC.cglasso](#), [summary.cglasso](#), [to_graph](#), [plot.cglasso2igraph](#) and [ShowStructure](#).

Examples

```
set.seed(123)
# Y ~ N(XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.5, probna = 0.05)
out <- cglasso(. ~ ., data = Z)

# MLE of the censored Gaussian graphical model identified by 'BIC'
out.mle <- cggm(out, GoF = BIC)
out.mle

# accessor functions
coef(out.mle, drop = TRUE)
fitted(out.mle, drop = TRUE)
residuals(out.mle, type = "working", drop = TRUE)
impute(out.mle, type = "both")

# goodness-of-fit functions
AIC(out.mle)
BIC(out.mle)
summary(out.mle)

# network analysis
out.graph <- plot(out.mle)
out.graph
```

cglasso

*Conditional Graphical Lasso Estimator***Description**

'cglasso' fits the conditional graphical lasso model to datasets with censored and/or missing values.

Usage

```
cglasso(formula, data, subset, contrasts = NULL, diagonal = FALSE,
         weights.B = NULL, weights.Tht = NULL, nlambda, lambda.min.ratio,
         lambda, nrho, rho.min.ratio, rho, maxit.em = 1.0E+4, thr.em = 1.0E-3,
         maxit.bcd = 1.0E+5, thr.bcd = 1.0E-4, trace = 0L)
```

Arguments

formula	an object of class 'formula' (or one that can be coerced to that class): a symbolic description of the model to be fitted.
data	an R object of S3 class 'datacggm', that is, the output of the function <code>datacggm</code> . See section 'Description' for more details.
subset	an optional vector specifying a subset of observations to be used in the fitting process.
contrasts	an optional list. See the <code>contrasts.arg</code> of <code>model.matrix.default</code> .
diagonal	logical. Should diagonal entries of the concentration matrix be penalized? Default is 'diagonal = FALSE'.
weights.B	an optional $q \times p$ dimensional matrix of non-negative weights used to penalize the regression coefficients (intercepts are unpenalized). This matrix can be also used to specify the unpenalized regression coefficients ('weights.B[i, j] = 0') or the structural zeros ('weights.B[i, j] = +Inf'). By default, all weights are set equal to 1, meaning that the penalized regression coefficients are unweighted.
weights.Tht	an optional symmetric matrix of non-negative weights used to penalize the partial regression coefficients. This matrix can be used to specify the unpenalized partial correlation coefficients ('weights.Tht[i, j] = 0') or the structural zeros in the precision matrix ('weights.Tht[i, j] = +Inf'). By default, the off-diagonal entries of the matrix 'weights.Tht' are set equal to 1, meaning that the partial correlation coefficients are unweighted, whereas the diagonal entries are equal to 0, that is the diagonal entries of the precision matrix are unpenalized.
nlambda	integer. The number of λ -values used to penalize the regression coefficients. By default 'nlambda = 10'.
lambda.min.ratio	the smallest λ -value is defined as a fraction of 'lambda.max' (i.e., the smallest λ -value for which all the estimated regression coefficients are equal to zero). The default depends on the sample size 'n' relative to the number of predictors 'q'. If 'q < n', default is '1.0E-6' otherwise the value '1.0E-2' is used as default.

A very small value of ‘lambda.min.ratio’ will lead to a saturated fitted model in the ‘ $q < n$ ’ case.

lambda	an optional user-supplied decreasing sequence of λ -values. By default cglasso computes a sequence of λ -values using nlambda and lambda.min.ratio. If lambda is supplied then nlambda and lambda.min.ratio are overwritten. WARNING: use with care and avoid supplying a single λ -value; supply instead a decreasing sequence.
nrho	integer. The number of ρ -values used to penalize the partial correlation coefficients. By default ‘nrho = 10’.
rho.min.ratio	the smallest ρ -value is defined as a fraction of ‘rho.max’ (i.e., the smallest ρ -value for which all the estimated partial correlation coefficients are equal to zero). The default depends on the sample size ‘ n ’ relative to the number of response variables ‘ p ’. If ‘ $p < n$ ’, the default is ‘1.0E-6’ otherwise the value ‘1.0E-2’ is used as default. A very small value of ‘rho.min.ratio’ will lead to a saturated fitted model in the ‘ $p < n$ ’ case.
rho	an optional user supplied decreasing sequence of ρ -values. By default cglasso computes a sequence of ρ -values using nrho and rho.min.ratio. If rho is supplied then nrho and rho.min.ratio are overwritten. WARNING: use with care and avoid supplying a single ρ -value; supply instead a decreasing sequence.
maxit.em	maximum number of iterations of the EM algorithm. Default is 1.0E+4.
thr.em	threshold for the convergence of the EM algorithm. Default value is 1.0E-4.
maxit.bcd	maximum number of iterations of the glasso algorithm. Default is 1.0E+5.
thr.bcd	threshold for the convergence of the glasso algorithm. Default is 1.0E-4.
trace	integer for printing information out as iterations proceed: trace = 0 no information is printed out; trace = 1 minimal information is printed on screen; trace = 2 detailed information is printed on screen.

Details

cglasso is the main model-fitting function and can be used to fit a broad range of extensions of the glasso estimator (Friedman *and other*, 2008). It is specifically proposed to study datasets with censored and/or missing response values. To help the user, the cglasso function has been designed to automatically select the most suitable extension by using the information stored in the ‘[datacggm](#)’ object passed through Z.

Below we sum up the available extensions:

- if only left/right-censored are observed in the response matrix Y (without missing values) and no predictor matrix is stored in Z, then cglasso computes the censored glasso estimator (Augugliaro *and other*, 2020a);
- if only left/right-censored are observed in the response matrix Y (without missing values) and a predictor matrix X is stored in Z, then cglasso computes the conditional censored glasso estimator (Augugliaro *and other*, 2020b);
- if only missing values are stored in the response matrix Y (without censored values), then cglasso computes the missglasso estimator (Stadler *and other*, 2012);
- starting with version 2.0.0, cglasso can also handle datasets with both missing and censored response values.

See section ‘Examples’ for some example.

The model-fitting function `cglasso` returns an R object of S3 class ‘`cglasso`’ for which there are available a set of accessor functions, a set of functions designed to evaluate the goodness-of-fit of the fitted models and, finally, a set of functions developed to analyze the selected network. The function `ShowStructure` can be used to show the structure of the package.

The accessor functions `coef.cglasso`, `fitted.cglasso`, `residuals.cglasso`, `predict.cglasso` and `impute` can be used to extract various useful features of the object fitted by `cglasso`.

For an R object returned by `cglasso`, the functions `AIC.cglasso` and `BIC.cglasso` can be used to evaluate the goodness-of-fit of the fitted models. Usually, these functions are used together with the function `summary.cglasso`, which gives more information about the sequence of fitted models. The plotting function `plot.GoF` can be used to graphically identify the optimal pair of the tuning parameters.

Given a pair of the tuning parameters, the functions `to_graph` and `plot.cglasso2igraph` can be used to analyze and show the selected network. Finally, the function `cggm` can be used to produce post-hoc maximum likelihood refitting of the selected graphical model.

Value

`cglasso` returns an object of S3 class “`cglasso`”, i.e., a named list containing the following components:

<code>call</code>	the call that produced this object.
<code>Yipt</code>	an array of dimension ‘ $n \times p \times n\lambda \times nrho$ ’ storing the ‘working response matrices’, that is, <code>Yipt[, , i, j]</code> is used as response matrix to compute the multilasso estimator (Augugliaro <i>and other</i> , 2020b). The accessor function ‘ <code>impute</code> ’ can be used to extract the desired imputed matrix.
<code>B</code>	an array of dimension ‘ $(q + 1) \times p \times n\lambda \times nrho$ ’ storing the penalized estimate of the regression coefficient matrices. The accessor function ‘ <code>coef.cglasso</code> ’ can be used to extract the desired estimates.
<code>mu</code>	an array of dimension ‘ $n \times p \times n\lambda \times nrho$ ’ storing the fitted values. The accessor function ‘ <code>fitted.cglasso</code> ’ can be used to extract the desired fitted values.
<code>R</code>	an array of dimension ‘ $n \times p \times n\lambda \times nrho$ ’ storing the ‘working residuals’, that is, <code>R[, , i, j]</code> is defined as the difference between <code>Yipt[, , i, j]</code> and <code>mu[, , i, j]</code> . The accessor function ‘ <code>residuals.cglasso</code> ’ can be used to extract the desired residual matrix.
<code>S</code>	an array of dimension ‘ $p \times p \times n\lambda \times nrho$ ’ storing the ‘working empirical covariance matrices’, that is, ‘ <code>S[, , i, j]</code> ’ is used to compute the glasso estimator.
<code>Sgm</code>	an array of dimension ‘ $p \times p \times n\lambda \times nrho$ ’ storing the estimated covariance matrices. The accessor function ‘ <code>coef</code> ’ can be used to extract the desired estimates.
<code>Tht</code>	an array of dimension ‘ $p \times p \times n\lambda \times nrho$ ’ storing the estimated precision matrices. The accessor function ‘ <code>coef</code> ’ can be used to extract the desired estimates.

dfB	a matrix of dimension ' $(\rho + 1) \times n\lambda \times nrho$ ' storing the number of estimated non-zero regression coefficients. Only for internal purpose.
dfTht	a matrix of dimension ' $n\lambda \times nrho$ ' storing the number of estimated non-zero (off-diagonal) partial correlation coefficients. Only for internal purpose.
InfoStructure	a named list whose elements contain information about the estimated networks. Only for internal purpose.
nit	an array of dimension ' $2 \times n\lambda \times nrho$ ' storing the number of EM steps.
Z	the 'datacggm' object used to compute the censored graphical lasso estimator.
diagonal	the flag used to specify if the diagonal entries of the precision matrix are penalized.
weights.B	the matrix of non-negative weights used for the regression coefficients.
weights.Tht	the matrix of non-negative weights used for the precision matrix.
nlambda	the number of λ -values used.
lambda.min.ratio	the value used to compute the smallest λ -value.
lambda	the sequence of λ -values used to fit the model.
nrho	the number of ρ -values used.
rho.min.ratio	the value used to compute the smallest ρ -value.
rho	the sequence of ρ -values used to fit the model.
model	a description of the fitted model.
maxit.em	maximum number of iterations of the EM algorithm.
thr.em	threshold for the convergence of the EM algorithm.
maxit.bcd	maximum number of iterations of the glasso algorithm.
thr.bcd	threshold for the convergence of the glasso algorithm.
conv	a description of the error that has occurred.
subrout	the name of the Fortran subroutine where the error has occurred (for internal debug only).
trace	the integer used for printing information on screen.
nobs	the sample size
nresp	the number of response variables used to fit the model.
npred	the number of predictors used to fit the model.

Author(s)

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References

- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020a) <[doi:10.1093/biostatistics/kxy043](https://doi.org/10.1093/biostatistics/kxy043)>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.
- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020b) <[doi:10.1007/s11222020099457](https://doi.org/10.1007/s11222020099457)>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
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- Stadler, N. and Buhlmann, P. (2012) <[doi:10.1007/s1122201092197](https://doi.org/10.1007/s1122201092197)>. Missing values: sparse inverse covariance estimation and an extension to sparse regression. *Statistics and Computing* **22**, 219–235.

See Also

[datacggm](#), [coef.cglasso](#), [fitted.cglasso](#), [residuals.cglasso](#), [predict.cglasso](#), [impute](#), [AIC.cglasso](#), [BIC.cglasso](#), [summary.cglasso](#), [select.cglasso](#), [plot.Gof](#), [to_graph](#), [plot.cglasso2igraph](#), [cggm](#) and [ShowStructure](#).

Examples

```
set.seed(123)

# Model 1: censored glasso estimator (Augugliaro \emph{and other}, 2020a)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out

# Model 2: conditional censored glasso estimator (Augugliaro \emph{and other}, 2020b)
# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out

# Model 3: missglasso estimator (Stadler \emph{and other}, 2012)
# Y ~ N(b0 + XB, Sigma) and probability of missing-at-random values equal to 0.05
n <- 1000L
p <- 3L
q <- 2L
```

```

b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probna = 0.05)
out <- cglasso(. ~ ., data = Z)
out

# Model 4: mixed estimator
# Y ~ N(b0 + XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 1000L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, X = X, b0 = b0, B = B, Sigma = Sigma, prob1 = 0.05, prob2 = 0.05,
           probna = 0.05)
out <- cglasso(. ~ ., data = Z)
out

```

coef

*Extract Model Coefficients***Description**

The accessor function `coef` extracts model coefficients from an R object inheriting class `'cglasso'`.

Usage

```

## S3 method for class 'cglasso'
coef(object, type = c("all", "B", "Sigma", "Theta"), lambda.id, rho.id,
     drop = TRUE, ...)

```

Arguments

<code>object</code>	an R object inheriting class <code>'cglasso'</code> , that is, the output of the model-fitting functions <code>cglasso</code> and <code>cggm</code> .
<code>type</code>	a description of the desired estimates.
<code>lambda.id</code>	an optional vector of integers used to specify the λ -values.
<code>rho.id</code>	an optional vector of integers used to specify the ρ -values.
<code>drop</code>	logical. Dimensions of the required objects can only be dropped if their extent is one.
<code>...</code>	further arguments passed to or from other methods.

Value

Coefficients extracted from ‘object’ are returned. By default, a named list storing all the estimated parameters is returned.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

Model-fitting functions [cglasso](#), [cggm](#) and the accessor functions [fitted.cglasso](#), [residuals.cglasso](#), [predict.cglasso](#) and [impute](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
coef(out, type = "Theta", rho.id = 1:4)
coef(out, type = "Theta", rho.id = 3, drop = TRUE)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
coef(out, type = "B", lambda.id = 3, rho.id = 1:4)
coef(out, type = "B", lambda.id = 3, rho.id = 3, drop = TRUE)
```

ColMeans + ColMeans *Calculate Column Means and Vars of a “datacggm” Object*

Description

Retrieve column means and column variances of a “datacggm” object.

Usage

```
ColMeans(x)
ColVars(x)
```

Arguments

`x` an object of class 'datacggm'.

Details

For an R object `x` of class 'datacggm', `ColMeans` (`ColVars`) retrieves the column means (variances) of the matrices obtained by `getMatrix(x, "Y")` and `getMatrix(x, "X")`. For the response variables, marginal means and variances are estimated using a EM-algorithm under the assumption that the p response variables are marginally normally distributed (see also Details section in [datacggm](#)). For the numeric predictor variables, marginal means and variances are computed by [mean](#) and [var](#), whereas, for categorical data, `ColMeans` (`ColVars`) retrieves the statistical mode and the Gini-Simpson Index, respectively.

Value

`ColMeans` (`ColVars`) returns a named list with the columns means (variances).

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacggm](#), [rcggm](#), [qqcnorm](#) and [hist.datacggm](#).

Examples

```
set.seed(123)
n <- 1000L
p <- 3L
b0 <- rep(0, p)
Z <- rcggm(n = n, b0 = b0, probl = 0.05, probr = 0.05)
ColMeans(Z)
ColVars(Z)

n <- 1000L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
Z <- rcggm(n = n, b0 = b0, X = X, B = B, probl = 0.05, probr = 0.05)
ColMeans(Z)
ColVars(Z)
```

datacggm	<i>Create a Dataset from a Conditional Gaussian Graphical Model with Censored and/or Missing Values</i>
----------	---

Description

‘The datacggm’ function is used to create a dataset from a conditional Gaussian graphical model with censored and/or missing values.

Usage

```
datacggm(Y, lo = -Inf, up = +Inf, X = NULL, control = list(maxit = 1.0E+4,
  thr = 1.0E-4))
```

Arguments

Y	a $(n \times p)$ -dimensional matrix; each row is an observation from a conditional Gaussian graphical model with censoring vectors lo and up. Missing-at-random values are recorded as ‘NA’.
lo	the lower censoring vector; lo[j] is used to specify the lower censoring value for the random variable Y_j .
up	the upper censoring vector; up[j] is used to specify the upper censoring value for the random variable Y_j .
X	an optional $(n \times q)$ -dimensional data frame of predictors. If missing (default), a dataset from a Gaussian graphical model is returned otherwise a dataset from a conditional Gaussian graphical model is returned.
control	a named list used to pass the arguments to the EM algorithm (see below for more details). The components are: <ul style="list-style-type: none"> • maxit: maximum number of iterations. Default is 1.0E+4. • thr: threshold for the convergence. Default value is 1.0E-4.

Details

The function ‘datacggm’ returns an R object of class ‘datacggm’, that is a named list containing the elements needed to fit a conditional graphical LASSO (cglasso) model to datasets with censored and/or missing values.

A set of specific method functions are developed to describe data with censored/missing values. For example, the method function ‘print.datacggm’ prints out the left and right-censored values using the following rules: a right-censored value is labeled adding the symbol ‘+’ at the end of the value, whereas the symbol ‘-’ is used for the left-censored values (see examples below). The summary statistics can be obtained using the method function ‘summary.datacggm’. The matrices Y and X are extracted from a datacggm object using the function ‘getMatrix’.

For each column of the matrix ‘Y’, mean and variance are estimated using a standard EM-algorithm based on the assumption of a Gaussian distribution. ‘maxit’ and ‘thr’ are used to set the number of iterations and the threshold for convergence, respectively. Marginal means and variances can be

extracted using the accessor functions ‘`ColMeans`’ and ‘`ColVars`’, respectively. Furthermore, the plotting functions ‘`hist.datacggm`’ and ‘`qqcnorm`’ can be used to inspect the marginal distribution of each column of the matrix ‘`Y`’.

The status indicator matrix, denoted by `R`, can be extracted by using the function `event`. The entries of this matrix specify the status of an observation using the following code:

- ‘`R[i, j] = 0`’ means that the y_{ij} is inside the open interval $(lo[j], up[j])$;
- ‘`R[i, j] = -1`’ means that the y_{ij} is a left-censored value;
- ‘`R[i, j] = +1`’ means that the y_{ij} is a right-censored value;
- ‘`R[i, j] = +9`’ means that the y_{ij} is a missing value.

See below for the other functions related to an object of class ‘`datacggm`’.

Value

‘`datacggm`’ returns an R object of S3 class “`datacggm`”, that is, a nested named list containing the following components:

<code>Y</code>	the $(n \times p)$ -dimensional matrix <code>Y</code> .
<code>X</code>	the $(n \times q)$ -dimensional data frame <code>X</code> .
<code>Info</code>	<ul style="list-style-type: none"> • <code>lo</code>: the lower censoring vector; • <code>up</code>: the upper censoring vector; • <code>R</code>: the status indicator matrix encoding the censored/missing values (mainly for internal purposes); • <code>order</code>: an integer vector used for the ordering of the matrices <code>Y</code> and <code>X</code> (for internal purposes only); • <code>Pattern</code>: a matrix encoding the information about the the patterns of censored/missing values (for internal purposes only); • <code>ym</code>: the estimated marginal means of the random variables Y_j; • <code>yv</code>: the estimated marginal variances of the random variables Y_j; • <code>n</code>: the sample size; • <code>p</code>: the number of response variables; • <code>q</code>: the number of columns of the data frame <code>X</code>.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020a) <[doi:10.1007/s11222020099457](https://doi.org/10.1007/s11222020099457)>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020b) <[doi:10.1093/biostatistics/kxy043](https://doi.org/10.1093/biostatistics/kxy043)>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

Related to the R objects of class “datacggm” there are the accessor functions, [rowNames](#), [colNames](#), [getMatrix](#), [ColMeans](#), [ColVars](#), [upper](#), [lower](#), [event](#), [qqcnorm](#) and the method functions [is.datacggm](#), [dim.datacggm](#), [summary.datacggm](#) and [hist.datacggm](#). The function [rcggm](#) can be used to simulate a dataset from a conditional Gaussian graphical model whereas [cglasso](#) is the model fitting function devoted to the l1-penalized censored Gaussian graphical model.

Examples

```
set.seed(123)

# a dataset from a right-censored Gaussian graphical model
n <- 100L
p <- 3L
Y <- matrix(rnorm(n * p), n, p)
up <- 1
Y[Y >= up] <- up
Z <- datacggm(Y = Y, up = up)
Z

# a dataset from a conditional censored Gaussian graphical model
n <- 100L
p <- 3L
q <- 2
Y <- matrix(rnorm(n * p), n, p)
up <- 1
lo <- -1
Y[Y >= up] <- up
Y[Y <= lo] <- lo
X <- matrix(rnorm(n * q), n, q)
Z <- datacggm(Y = Y, lo = lo, up = up, X = X)
Z

# a dataset from a conditional censored Gaussian graphical model
# and with missing-at-random values
n <- 100L
p <- 3L
q <- 2
Y <- matrix(rnorm(n * p), n, p)
NA.id <- matrix(rbinom(n * p, 1L, 0.01), n, p)
Y[NA.id == 1L] <- NA
up <- 1
lo <- -1
Y[Y >= up] <- up
Y[Y <= lo] <- lo
X <- matrix(rnorm(n * q), n, q)
Z <- datacggm(Y = Y, lo = lo, up = up, X = X)
Z
```

dim.datacggm	<i>Dimensions of a “datacggm” Object</i>
--------------	--

Description

Retrieve the dimension of an R object of class “datacggm”, that is, the sample size (n), the number of response variables (p) and the number of predictors (q).

Usage

```
## S3 method for class 'datacggm'  
dim(x)
```

Arguments

x an R object of class ‘datacggm’.

Details

For an R object of class ‘datacggm’, dim retrieves a list with elements named Y and X. The first component is the “dim” attribute of the matrix Y whereas the second component is the “dim” attribute of the matrix X. If X is missing, the second component is NULL.

Value

For an object of class ‘datacggm’, dim retrieves a named list with components Y and X which are the “dim” attribute of the two matrices, respectively. If X is missing then the last component is NULL.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacggm](#), [rcggm](#), [nobs](#), [nresp](#) and [npred](#).

Examples

```
set.seed(123)  
# a dataset from a censored Gaussian graphical model  
n <- 100L  
p <- 3L  
b0 <- rep(0, p)  
Z <- rcggm(n = n, b0 = b0, probl = 0.05, probr = 0.05)  
dim(Z)  
  
# a dataset from a conditional censored Gaussian graphical model  
n <- 100L  
p <- 3L
```

```
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
Z <- rcggm(n = n, b0 = b0, X = X, B = B, probl = 0.05, probr = 0.05)
dim(Z)
```

dimnames.datacggm *Dimnames of a “datacggm” Object*

Description

Retrieve or set the dimnames of a “datacggm” object.

Usage

```
## S3 method for class 'datacggm'
dimnames(x)

## S3 replacement method for class 'datacggm'
dimnames(x) <- value
```

Arguments

x an object of class ‘datacggm’.

value a nested list with names ‘X’ and ‘Y’. See below for mode details.

Details

For an R object ‘x’ of class ‘datacggm’, `dimnames` retrieves or sets the `dimnames` attribute for the matrices Y and X (see ‘[datacggm](#)’ for more details). When setting the `dimnames` attribute, `value$Y` can be NULL (which is not stored) or a list of length two. In the last case, `value$Y` is passed to the method function `dimnames` for setting the `dimnames` attributes of the matrix retrieved by `getMatrix(x, "Y")`. In the same way, `value$X` can be used for setting the `dimnames` attributes of the matrix retrieved by `getMatrix(x, "X")`.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacggm](#), [rcggm](#), [getMatrix](#), [rowNames](#) and [colNames](#).

Examples

```

set.seed(123)
# a dataset from a censored Gaussian graphical model
n <- 100L
p <- 3L
b0 <- rep(0, p)
Z <- rcggm(n = n, b0 = b0, probl = 0.05, probr = 0.05)
dimnames(Z)

dimnames(Z) <- list(Y = list(paste0("i", seq_len(n)), paste("Y", 1:p, sep = ":")),
dimnames(Z)

# the same as
# dimnames(Z)$Y <- list(paste0("i", seq_len(n)), paste("Y", 1:p, sep = ":"))
# dimnames(Z)

# a dataset from a conditional censored Gaussian graphical model
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
Z <- rcggm(n = n, b0 = b0, X = X, B = B, probl = 0.05, probr = 0.05)
dimnames(Z)

dimnames(Z) <- list(Y = list(NULL, paste("Y", 1:p, sep = ":")),
                    X = list(NULL, paste("X", 1:q, sep = ":")))
dimnames(Z)

# the same as
# dimnames(Z)$Y <- list(NULL, paste("Y", 1:p, sep = ":"))
# dimnames(Z)$X <- list(NULL, paste("X", 1:q, sep = ":"))
# dimnames(Z)

```

event

Status Indicator Matrix from a 'datacggm' Object

Description

The 'event' function retrieves the status indicator matrix from an object of class 'datacggm'.

Usage

```
event(x, ordered = FALSE)
```

Arguments

<code>x</code>	an object of class ‘datacggm’.
<code>ordered</code>	logical value used to specify if the rows of the status indicator matrix should be ordered according to the patterns of censored/missing values. Default <code>ordered = FALSE</code> .

Details

The ‘event’ function is used to retrieve the status indicator matrix, denoted by R , from an object of class ‘datacggm’. The entries of the matrix are used to specify the status of the response variable:

- ‘ $R[i, j] = 0$ ’ means that y_{ij} is inside the open interval $(lo[j], up[j])$;
- ‘ $R[i, j] = -1$ ’ means that y_{ij} is a left-censored value;
- ‘ $R[i, j] = +1$ ’ means that y_{ij} is a right-censored value;
- ‘ $R[i, j] = +9$ ’ means that y_{ij} is a missing value.

Value

event returns a $(n \times p)$ -dimensional matrix.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

Augugliaro, L., Sottile, G., and Vinciotti, V. (2020) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.

Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#) and [rcggm](#).

Examples

```
set.seed(123)

# Y ~ N(b0 + XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random equals to 0.05
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
```

```

Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05,
          probr = 0.05, probna = 0.05)

# status indicator matrix
event(Z)

# in this case the status indicator matrix is returned with
# rows ordered according to the patterns of missing data
event(Z, ordered = TRUE)

```

Example
Simulated data for the cglasso vignette

Description

Simulated data, used to demonstrate the features of `datacggm` class. It contains two simulated datasets that mimic the structural characteristics of the two real datasets MKMEP and MM.

1. `MKMEP.Sim`: is a response matrix containing 50 samples of 10 genes;
2. `MM.Sim`: is a list containing two components;
 - `Y`: is the response matrix containing 50 measurements of 10 miRNAs
 - `X`: is the predictor matrix containing 50 measurements of 5 variables (four numerical and one categorical)

In both datasets, the response matrices are right-censored with an upper limit of detection fixed to 40.

Usage

```
data("Example")
```

See Also

[cglasso](#), [to_graph](#), and the method functions [summary](#), [coef](#), [plot](#), [AIC.cglasso](#), [BIC.cglasso](#), [MKMEP](#) and [MM](#).

Examples

```

data("Example")

MM.Sim <- datacggm(Y = MM.Sim$Y, up = 40, X = MM.Sim$X)
ColMeans(MM.Sim)
ColVars(MM.Sim)
summary(MM.Sim)

MKMEP.Sim <- datacggm(Y = MKMEP.Sim, up = 40)
ColMeans(MKMEP.Sim)
ColVars(MKMEP.Sim)
summary(MKMEP.Sim)

```

fitted *Extract Model Fitted Values*

Description

The accessor function `fitted` extracts model fitted values from an R object inheriting class `'cglasso'`.

Usage

```
## S3 method for class 'cglasso'
fitted(object, lambda.id, rho.id, drop = TRUE, ...)
```

Arguments

<code>object</code>	an R object inheriting class <code>'cglasso'</code> , that is, the output of the model-fitting functions <code>cglasso</code> or <code>cggm</code> .
<code>lambda.id</code>	a vector of integers used to specify the λ -values.
<code>rho.id</code>	a vector of integers used to specify the ρ -values.
<code>drop</code>	logical. Dimensions can only be dropped if their extent is one.
<code>...</code>	further arguments passed to or from other methods.

Value

Fitted values extracted from `'object'` are returned.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

Model-fitting functions `cglasso`, `cggm` and the accessor functions `coef.cglasso`, `residuals.cglasso`, `predict.cglasso` and `impute`.

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
fitted(out, rho.id = 3L, drop = TRUE)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
```

```

p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
fitted(out, lambda.id = 3L, rho.id = 3L, drop = TRUE)

```

getGraph

Retrieve Graphs from a 'cglasso2igraph' Object

Description

'getGraph' retrieves graphs from an R object of class 'cglasso2igraph'.

Usage

```
getGraph(x, type = c("Gyy", "Gxy", "both"))
```

Arguments

x an object of class 'cglasso2igraph' (see also [to_graph](#)).
type a description of the required graph. Default is 'Gyy'.

Value

getGraph retrieves an R object of class 'igraph' representing the graph required by the argument type.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[to_graph](#), [is.cglasso2igraph](#) and [plot.cglasso2igraph](#).

Examples

```

set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)

```

```

out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out)
getGraph(out.graph)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out, lambda.id = 3, rho.id = 3, weighted = TRUE)
getGraph(out.graph, type = "Gyy")
getGraph(out.graph, type = "Gxy")
getGraph(out.graph, type = "both")

```

getMatrix

Retrieve Matrices 'Y' and 'X' from a 'datacggm' Object

Description

'getMatrix' retrieves matrices 'Y' and/or 'X' from an object of class 'datacggm'.

Usage

```
getMatrix(x, name = c("Y", "X", "both"), ordered = FALSE)
```

Arguments

x	an object of class 'datacggm'.
name	the name of the required matrix.
ordered	logical value used to specify if the required matrix should be retrieved with rows ordered according to the patterns of censored values. See below for some example.

Value

getMatrix retrieves the matrix specified by 'name' and with row ordering specified by 'ordered'. A named list returned if name is "both".

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

Augugliaro, L., Sottile, G., and Vinciotti, V. (2020) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.

Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#), [rcggm](#).

Examples

```
set.seed(123)

# a dataset from a conditional censored Gaussian graphical model
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
Z <- rcggm(n = n, b0 = b0, X = X, B = B, probl = 0.05, probr = 0.05)
getMatrix(Z, name = "Y")

# in the following examples 'Y' and 'X' is returned with rows ordered
# according to the patterns of censored data
getMatrix(Z, name = "Y", ordered = TRUE)

getMatrix(Z, name = "X")
getMatrix(Z, name = "X", ordered = TRUE)

getMatrix(Z, name = "both")
getMatrix(Z, name = "both", ordered = TRUE)
```

hist.datacggm

Histogram for a datacggm Object

Description

Creates histograms with a normal density as background and two segments corresponding to left and/or right censored values (if any) in a conditional censored Gaussian graphical model.

Usage

```
## S3 method for class 'datacggm'
hist(x, breaks = "Sturges", include.lowest = TRUE, right = TRUE, nclass = NULL,
     which, max.hist = 1L, save.hist = FALSE, grdev = pdf, grdev.arg,
     polygon.col = adjustcolor("grey", alpha.f = 0.25), polygon.border = NA,
```

```
segments.lwd = 4L, segments.lty = 2L, segments.col = "gray40",
points.pch = c(4L, 1L), points.cex = 1.8, points.col = rep("black", 2L),
legend = TRUE, ...)
```

Arguments

<code>x</code>	an object of class 'datacggm'.
<code>breaks</code>	one of: <ul style="list-style-type: none"> • a vector giving the breakpoints between histogram cells, • a function to compute the vector of breakpoints, • a single number giving the number of cells for the histogram, • a character string naming an algorithm to compute the number of cells, • a function to compute the number of cells. See hist for more details.
<code>include.lowest</code>	logical; if TRUE, an <code>x[i]</code> equal to the <code>breaks</code> value will be included in the first (or last, for <code>right = FALSE</code>) bar. This will be ignored (with a warning) unless <code>breaks</code> is a vector.
<code>right</code>	logical; if TRUE, the histogram cells are right-closed (left open) intervals.
<code>nclass</code>	numeric (integer). For S(-PLUS) compatibility only, <code>nclass</code> is equivalent to <code>breaks</code> for a scalar or character argument.
<code>which</code>	a vector of integers used to specify the response variables for which the histogram is required.
<code>max.hist</code>	the maximum number of histograms drawn in a single figure.
<code>save.hist</code>	a logical variable or a string specifying the path of the directory where plots will be saved. Letting ' <code>save.hist = TRUE</code> ', the required plots will be saved as external files inside the current working directory. User can save these files on a specific working directory passing the absolute path through the argument <code>save.hist</code> . On exit, working directory will be always set the previous one.
<code>grdev</code>	the graphics device used to save the required histograms on external files. See ' device ' for more details.
<code>grdev.arg</code>	additional parameters passed to the graphics device specified by ' <code>grdev</code> '.
<code>polygon.col</code>	graphical parameter; the colour for filling the area underlying the Gaussian distribution.
<code>polygon.border</code>	graphical parameter; the colour of the border of the Gaussian distribution.
<code>segments.lwd</code>	graphical parameter; the line width used in plotting the segments associated to the symbols specified by ' <code>points.pch</code> '.
<code>segments.lty</code>	graphical parameter; the line type used in plotting the segments associated to the symbols specified by ' <code>points.pch</code> '.
<code>segments.col</code>	graphical parameter; the colour used in plotting the segments associated to the symbols specified by ' <code>points.pch</code> '.
<code>points.pch</code>	graphical parameter; a pair of integers specifying the symbols used to plot the proportion of censored response variables (default ' <code>points.pch = 4L</code> ') and the probability to observe a censored value (default ' <code>points.pch = 1L</code> '), respectively.

<code>points.cex</code>	graphical parameter; a numerical value giving the amount by which the symbols, specified by <code>'points.pch'</code> , are magnified relative to the default.
<code>points.col</code>	graphical parameter; the colours used to plot the symbols specified by <code>'points.pch'</code> .
<code>legend</code>	logical; if <code>'TRUE'</code> legend is added to the plot.
<code>...</code>	additional graphical parameters passed to <code>'plot.histogram'</code> .

Details

For each response variable, the method function `'hist.datacggm'` plots a histogram using only the observed values. Densities, plotted on the y-axis, are computed in such a way that the sum of all densities plus the proportion of censored values is equal to one.

To evaluate the distributional assumption underlying the censored Gaussian distribution, on each histogram the area underlying the Gaussian density function is also shown in the background (marginal parameters are estimated as described in `'datacggm'` and `'ColMeans'`). Furthermore, on each plot, the proportions of left/right censored values are graphically compared with the corresponding Gaussian tail probabilities. If the assumption about the censoring mechanism is satisfied, then the proportion of censored values and the tail probability are approximately equal to each other.

Author(s)

Gianluca Sottile (<gianluca.sottile@unipa.it>)

See Also

[datacggm](#), [rcggm](#), [ColMeans](#), [ColVars](#) and [qqcnorm](#).

Examples

```
set.seed(123)

# a dataset from a right-censored Gaussian graphical model
n <- 1000L
p <- 10L
Y <- matrix(rnorm(n * p), n, p)
up <- 1
Y[Y >= up] <- up
Z <- datacggm(Y = Y, up = up)
hist(Z, max.hist = 4L)
```

Description

Imputes multivariate missing and censored values.

Usage

```
impute(object, type = c("mar", "censored", "both"), lambda.new, rho.new)
```

Arguments

object	an R object inheriting class 'cglasso', that is, the output of the model-fitting functions cglasso and cggm .
type	a description of the imputation required (see section 'Description'). By default only the missing-at-random values are imputed.
lambda.new	value of the tuning parameter λ at which the imputations are required.
rho.new	value of the tuning parameter ρ at which the imputations are required.

Details

The `impute` function returns the response matrix with the imputed missing-at-random values ('type = "mar"'), the imputed censored values ('type = "censored"'), or both ('type = "both"').

If 'type = "mar"' then, for each row, the missing response values are replaced with the expected values of a multivariate normal distribution conditioned on the observed response values.

If 'type = "censored"' then, for each row, the censored response values are imputed using the expected values of a multivariate truncated normal distribution conditioned on the observed response values.

If 'type = "both"' then missing-at-random and censored values are imputed. In this case the returned matrix corresponds to the working response matrix computed during the E-Step.

Value

The `impute` function returns a response matrix with the required imputed data.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

Model-fitting functions [cglasso](#), [cggm](#) and the accessor functions [coef.cglasso](#), [fitted.cglasso](#), [residuals.cglasso](#) and [predict.cglasso](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05, probna = 0.05)
```

```

out <- cglasso(. ~ ., data = Z)
rho.new <- mean(out$rho)

# imputing missing values
Y.impute <- impute(out, type = "mar", rho.new = rho.new)

# imputing censored values
Y.impute <- impute(out, type = "censored", rho.new = rho.new)

# imputing missing and censored values
Y.impute <- impute(out, type = "both", rho.new = rho.new)

# Y ~ N(XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05,
          probna = 0.05)

out <- cglasso(. ~ ., data = Z)
lambda.new <- mean(out$lambda)
rho.new <- mean(out$rho)

# imputing missing values
Y.impute <- impute(out, type = "mar", lambda.new = lambda.new, rho.new = rho.new)

# imputing censored values
Y.impute <- impute(out, type = "censored", lambda.new = lambda.new, rho.new = rho.new)

# imputing missing and censored values
Y.impute <- impute(out, type = "both", lambda.new = lambda.new, rho.new = rho.new)

```

is.cglasso2igraph *Is an Object of Class 'cglasso2igraph'?*

Description

is.cglasso2igraph tests if its argument is an object of class 'cglasso2igraph'.

Usage

```
is.cglasso2igraph(x)
```

Arguments

x object to be tested.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[to_graph](#).

Examples

```
set.seed(123)
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out)
is.cglasso2igraph(out.graph )
```

is.datacggm

Is an Object of Class 'datacggm'?

Description

'is.datacggm' tests if its argument is an object of class 'datacggm'.

Usage

```
is.datacggm(x)
```

Arguments

x object to be tested.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020a) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020b) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#) and [rcggm](#)

Examples

```
set.seed(123)
n <- 100L
p <- 3L
b0 <- rep(0, p)
Z <- rcggm(n = n, b0 = b0, probl = 0.05, probr = 0.05)
is.datacggm(Z) # TRUE
```

lower + upper

Lower and Upper Limits from a “datacggm” Object

Description

Functions ‘lower’ and ‘upper’ retrieve the lower and upper censoring values from an object of class “datacggm”.

Usage

```
lower(x)
upper(x)
```

Arguments

x an object of class ‘datacggm’.

Details

For an R object x of class ‘datacggm’, lower (upper) retrieves the lower (upper) censoring values of the response variables.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacggm](#)

Examples

```

set.seed(123)
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))

Z <- rcggm(n = n, Sigma = Sigma, probr = 0.05)
lower(Z)
upper(Z)

Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05)
lower(Z)
upper(Z)

Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
lower(Z)
upper(Z)

```

 MKMEP

Megakaryocyte-Erythroid Progenitors

Description

In a study about the formation of blood cells, Psaila *and others* (2016) have recently identified three distinct sub-populations of cells, which are all derived from hematopoietic stem cells through cell differentiation. One of these sub-populations, denoted by MK-MEP, is a previously unknown, rare population of cells that are bipotent but primarily generate megakaryocytic progeny.

Multiplex RT-qPCR has been used to profile 63 genes and 48 single human MK-MEP cells. RT-qPCR data are typically right-censored with a limit of detection fixed by the manufacturer to 40. Raw data have been mean normalized using the method proposed in Pipelers *and others* (2017). See Section 5 in Augugliaro *and others* (2018) for more details.

‘MKMEP’ is a matrix containing a subset of the data available from Psaila *and others* (2016).

Usage

```
data("MKMEP")
```

References

- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.
- Pipelers, P., Clement, L., Vynck, M., Hellemans, J., Vandesompele, J. and Thas, O. (2017) <doi:10.1371/journal.pone.0182832>. A unified censored normal regression model for qPCR differential gene expression analysis. *PLoS One* **12**, e0182832.

Psaila, B., Barkas, N., Iskander, D., Roy, A., Anderson, S., Ashley, N., Caputo, V. S., Lichtenberg, J., Loaiza, S., Bodine, D. M. *and others*. (2016) <doi: 10.1186/s13059-016-0939-7> Single-cell profiling of human megakaryocyte-erythroid progenitors identifies distinct megakaryocyte and erythroid differentiation pathways. *Genome Biology* **17**, 83–102.

See Also

[cglasso](#), [to_graph](#), and the method functions [summary](#), [coef](#), [plot](#), [AIC.cglasso](#) and [BIC.cglasso](#).

Examples

```
data("MKMEP")
is.matrix(MKMEP)
Z <- datacggm(Y = MKMEP, up = 40)
str(Z)
is.datacggm(Z)
print(Z, width = 80L)
```

Description

MicroRNAs (miRNAs) are endogenous small non-coding RNAs, approximately 22 nucleotides in length, that play a regulatory role in gene expression by mediating mRNA cleavage or translation expression. Several studies have shown that a deregulation of the miRNAs can cause a disruption in the gene regulation mechanisms of the cell and that this might even lead to cancerous phenotypes.

Gutierrez *and others* (2010) investigated the expression level of a set of 141 miRNAs, measured by RT-qPCR technology on a sample of 64 patients with multiple myeloma (MM). Patients were selected to represent the most relevant recurrent genetic abnormalities in MM. RT-qPCR data are typically right-censored and, in this study, the upper limit of detection was fixed to 40 cycles.

Formally, MM is a list with containing a subset of the data studied in Gutierrez *and others* (2010). Its components are:

- Y: the response matrix containing the cycle-threshold of the measured 49 miRNAs;
- X: the predictor matrix containing the expression levels of 15 endogenous internal reference genes, called housekeeping, and a factor encoding the cytogenetic abnormalities.

Usage

```
data("MM")
```

References

- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.
- Gutierrez, N. Sarasquete, M., Misiewicz-Krzeminska, I., Delgado, M. De Las Rivas, J., Ticona, F.V., Ferminan, E., Martin-Jimenez, P., Chillon, C., Risueno, A., Hernandez, J.M., Garcia-Sanz, R., Gonzalez, M. and San Miguel, J.F. (2010) <doi:10.1038/leu.2009.274>. Deregulation of microRNA expression in the different genetic subtypes of multiple myeloma and correlation with gene expression profiling. *Leukemia* **24**(3), 629–637.

See Also

[cglasso](#), [to_graph](#), and the method functions [summary](#), [coef](#), [plot](#), [AIC.cglasso](#) and [BIC.cglasso](#).

Examples

```
data("MM")
Z <- datacggm(Y = MM$Y, X = MM$X, up = 40)
print(Z, width = 80L)
```

nobs + nresp + npred *Extract the Number of Observations/Responses/Predictors from a datacggm Object*

Description

Extract the number of observations, response variables and predictors from an object of class `datacggm`.

Usage

```
## S3 method for class 'datacggm'
nobs(object, ...)
## S3 method for class 'datacggm'
nresp(object, ...)
## S3 method for class 'datacggm'
npred(object, ...)
```

Arguments

`object` an R object of class `datacggm`.
`...` further arguments to be passed to methods.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacgmm](#), [rcgmm](#) and [dim.datacgmm](#).

Examples

```

set.seed(123)
n <- 100
p <- 3
q <- 2
b0 <- rep(1, p)
X <- matrix(rnorm(n * q), n, q)
B <- matrix(rnorm(q * p), q, p)
Sigma <- outer(1:p, 1:p, function(i, j) 0.3^abs(i - j))
probl <- 0.05
probr <- 0.05
probna <- 0.05

Z <- rcgmm(n = n, b0 = b0, Sigma = Sigma, probl = probl, probr = probr,
          probna = probna)
nobs(Z)
nresp(Z)
npred(Z)

Z <- rcgmm(b0 = b0, X = X, B = B, Sigma = Sigma, probl = probl, probr = probr,
          probna = probna)
nobs(Z)
nresp(Z)
npred(Z)

```

plot.cgmm

Plot Method for a 'cgmm' Object

Description

The `plot.cgmm` produces graphs from an R object of class 'cgmm'.

Usage

```

## S3 method for class 'cgmm'
plot(x, type, weighted = FALSE, simplify = TRUE, ...)

```

Arguments

<code>x</code>	an R object of class 'cgmm', that is, the output of the function cgmm .
<code>type</code>	a description of the required graph. Default depends on the type of fitted model, that is, <code>type = both</code> if a conditional censored graphical lasso estimator is fitted, otherwise <code>type</code> is equal to <code>Gyy</code> (see getGraph) for more details.
<code>weighted</code>	logical. Should weighted graphs be created? Default is <code>FALSE</code> .

simplify logical. Should isolated vertices be removed from the graph? Default is TRUE, i.e., isolated vertices are removed.

... additional graphical arguments passed to the functions [plot.igraph](#).

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[cggm](#), [to_graph](#), [getGraph](#) and [plot.cglasso2igraph](#).

Examples

```
set.seed(123)
# Y ~ N(XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.5,
          probna = 0.05)
out <- cglasso(. ~ ., data = Z)
out.mle <- cggm(out, lambda.id = 3L, rho.id = 3L)
plot(out.mle, type = "Gyy")
plot(out.mle, type = "Gxy")
plot(out.mle, type = "both")
```

plot.cglasso

Plot Method for 'cglasso' Object

Description

'The `plot.cglasso`' function produces plots to study the coefficient paths of fitted `cglasso` models.

Usage

```
## S3 method for class 'cglasso'
plot(x, what = c("Theta", "diag(Theta)", "b0", "B"),
     penalty = ifelse(x$nrho >= x$nlambda, "rho", "lambda"), given = NULL,
     GoF = AIC, add.labels, matplot.arg1, matplot.arg2, labels.arg, abline.arg,
     mtext.arg, save.plot, grdev = pdf, grdev.arg, digits = 4L, ...)
```

Arguments

x	an R object of class 'cglasso', that is, the output of the fitting function cglasso .
what	a character or a formula specifying the required plot. If what is a character, then it is used to specify the conditional coefficient path. Allowed descriptors are: 'Theta', the off-diagonal entries of the precision matrix, 'diag(Theta)', the diagonal entries of the precision matrix, 'b0', the intercepts of the conditional models, or equivalently the expected values of the response variables, and 'B', the regression coefficients. Optionally, 'what' can be a model formula (see sections Description and Examples for more details).
penalty	optional character argument used to specify the tuning parameter needed to plot the conditional coefficient path. Allowed descriptors are 'rho' and 'lambda'. This argument can be omitted if what is a model formula.
given	an optional vector of integers identifying the conditioning values of the second tuning parameter to be used for the coefficient profile plots of the parameters identified by the input 'what' across their corresponding tuning parameter. This argument is required only if 'what' is a character, otherwise it can be omitted.
GoF	a valid goodness-of-fit function, such as AIC.cglasso and BIC.cglasso , or an R object of class 'GoF'.
add.labels	logical value. Should labels be added to the plot?
matplot.arg1	a named list with the graphical parameters used to plot the paths of the estimates identified by 'GoF'.
matplot.arg2	a named list with the graphical parameters used to plot the paths of the remaining estimates.
labels.arg	a named list with the graphical parameters used to plot the labels.
abline.arg	a named list with the graphical parameters used to plot the line identifying the optimal tuning parameter value.
mtext.arg	a named list with the graphical parameters used to plot the text reported on the third axis.
save.plot	a logical variable or a string specifying the path of the directory where plots will be saved. Letting 'save.plot = TRUE', the required plots will be saved as external files inside the current working directory. User can save these files on a specific working directory passing the absolute path through the argument save.plot. On exit, working directory will be always set the previous one.
grdev	the graphics device used to save the required histograms on external files. See ' device ' for more details.
grdev.arg	additional parameters passed to the graphics device specified by 'grdev'.
digits	number of digits used to print the value of the second tuning parameter identified by 'given'.
...	further arguments passed to the chosen goodness-of-fit function, such as 'AIC' or 'BIC'.

Details

The function `plot.cglasso` produces the coefficient profile plot. The output depends both on the type of fitted model and on the setting of the three main arguments, `what`, `penalty` and `given`. Below we give more details.

If the model fitting function `cglasso` is used to fit an l_1 -penalized censored Gaussian graphical model (see first part in Section Example), then the user can specify only the main argument `what`, whereas `penalty` and `given` can be omitted. If `penalty` is specified, then it must be equal to `"rho"`. The main argument `what` is used to specify the estimator that is plotted on the y -axis. In this case, it must be equal to one of the following descriptors:

- `"Theta"`: the path of the estimated partial correlation coefficients is returned;
- `"diag(Theta)"`: the path of the diagonal entries of the estimated precision matrix is returned;
- `"b0"`: the path of the estimated expected values of the response variables is returned;

If a l_1 -penalized conditional censored Gaussian graphical model is fitted (see second part in Section Example), then all the main arguments can be used. In this case:

1. `what` must be equal to one of the following descriptors:
 - `"Theta"`: the path of the off-diagonal entries of the estimated precision matrix is returned;
 - `"diag(Theta)"`: the path of the estimated partial correlation coefficients is returned;
 - `"b0"`: the path of the estimated intercepts is returned;
 - `"B"`: the path of the estimated regression coefficients is returned for each response variable;
2. `penalty` is used to specify the tuning parameter that is reported on the x -axis and it must be equal to one of the following descriptors:
 - `"rho"`: the tuning parameter used to penalize the precision matrix;
 - `"lambda"`: the tuning parameter used to penalize the regression coefficient matrix;
3. `given` is an optional vector of integers used to specify the values of the second tuning parameter. For instance, letting `what = "Theta"`, `penalty = "rho"` and `given = 1` then `plot.cglasso` returns the plot of the estimated partial correlation coefficients versus the ρ -values and `given` the first value of the used λ -values. If `given` is left unspecified then a sequence of conditional profile plot is returned, a plot for each value of the second tuning parameter.

Optionally, the user can specify a conditional profile plot using a model formula with the following template:

```
'what ~ penalty | given'
```

For instance, the previous plot can be specified using the following model formula

```
'Theta ~ rho | 1'
```

In this case, the arguments `penalty` and `given` must be left unspecified whereas `what` is used for the model formula (see also examples below).

The optional argument `GoF` can be used to identify the non-zero estimates by the output of a goodness-of-fit function, such as `AIC.cglasso` and `BIC.cglasso`. In this case, a vertical red dashed line is used to identify the optimal value of the tuning parameter reported on the x -axis, whereas the path of the identified non-zero estimates is drawn using a solid black line; the remaining paths are drawn using gray dashed lines. This option is disabled if we let `'GoF = NULL'`, when all the paths are drawn using solid black lines.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

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- Friedman, J.H., Hastie, T., and Tibshirani, R. (2008) <doi:10.1093/biostatistics/kxm045>. Sparse inverse covariance estimation with the graphical lasso. *Biostatistics* **9**, 432–441.
- Stadler, N. and Buhlmann, P. (2012) <doi:10.1007/s1122201092197>. Missing values: sparse inverse covariance estimation and an extension to sparse regression. *Statistics and Computing* **22**, 219–235.

See Also

[cglasso](#), [AIC.cglasso](#) and [BIC.cglasso](#).

Examples

```
set.seed(123)

#####
# Model 1: censored glasso estimator (Augugliaro \emph{and other}, 2020a)
#  $Y \sim N(0, \text{Sigma})$  and probability of left/right censored values equal to 0.05
#####
n <- 100L
p <- 10L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)

# coefficient profile plot of the off-diagonal entries of the precision matrix
plot(out, what = "Theta")
# the same as
#plot(out, what = "Theta", penalty = "rho")

# the output of a goodness-of-fit function can be used to identify the
# non-zero estimates
plot(out, what = "Theta", GoF = BIC)

# letting 'GoF = NULL' the previous option is disabled
plot(out, what = "Theta", GoF = NULL)

# coefficient profile plot of the diagonal entries of the precision matrix
plot(out, what = "diag(Theta)")

# coefficient profile plot of the expected values of the response variables
```

```

plot(out, what = "b0")

#####
# Model 2: conditional censored glasso estimator (Augugliaro \emph{and other}, 2020b)
#  $Y \sim N(b_0 + XB, \text{Sigma})$  and probability of left/right censored values equal to 0.05
#####
n <- 100L
p <- 10L
q <- 5L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probr = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)

# conditional profile plot of the estimator partial correlation coefficients versus
# the used values of the tunin parameter rho and given the first lambda-value
plot(out, what = "Theta", penalty = "rho", given = 1L)
out$lambda[1L]

lambda.id <- c(2, 4)
plot(out, what = "Theta", penalty = "rho", given = lambda.id)
out$lambda[lambda.id]

# in this case a sequence of ten conditional profile plots is returned, tha is a
# plot for each lambda-value.
plot(out, what = "Theta", penalty = "rho")

# optionally, the user can specify the conditional profile plots using the model
# formula
plot(out, what = Theta ~ rho | lambda.id)
lambda.id

plot(out, what = Theta ~ rho)

# the output of a goodness-of-fit function can be used to identify the
# non-zero estimates
plot(out, what = Theta ~ rho | 10, GoF = BIC)

# letting 'GoF = NULL' the previous option is disabled
plot(out, what = Theta ~ rho | 10, GoF = NULL)

```

plot.cglasso2igraph *Plot Method for a cglasso2igraph Object*

Description

plot.cglasso2igraph produces graphs from an R object of class 'cglasso2igraph'.

Usage

```
## S3 method for class 'cglasso2igraph'
plot(x, type, ...)
```

Arguments

x an R object of class 'cglasso2igraph', that is, the output of the function [to_graph](#).
type a description of the required graph. Default is 'both' in a conditional glasso estimator or 'Gyy' in a glasso estimator (see [getGraph](#)).
... additional graphical arguments passed to the functions [plot.igraph](#).

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[to_graph](#) and [getGraph](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcgmm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out)
plot(out.graph, type = "Gyy")

out.graph <- to_graph(out, weighted = TRUE)
plot(out.graph, type = "Gyy")

# Y ~ N(b0 +XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcgmm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out, lambda.id = 3, rho.id = 3, weighted = TRUE)
plot(out.graph, type = "Gyy")
plot(out.graph, type = "Gxy")
plot(out.graph, type = "both")
```

plot.GoF *Plot for 'GoF' Object*

Description

'The plot.GoF' function produces plots to study the sequence of fitted models.

Usage

```
## S3 method for class 'GoF'
plot(x, add.line = TRUE, arg.line = list(lty = 2L, lwd = 2L, col = "red"),
     add.text = FALSE, arg.text = list(side = 3L), arg.points = list(pch = 2L),
     ...)
```

Arguments

x	an R object of class 'GoF', that is, the output of a goodness-of-fit function such as AIC.cglasso or BIC.cglasso .
add.line	logical; if 'add.line = TRUE' then a line is added to identify the optimal value of the tuning parameter.
arg.line	a named list of graphical parameters passed to the function abline (see also par).
add.text	logical; if 'add.text = TRUE' then a text is added to the line used to identify the optimal value of the tuning parameter.
arg.text	a list of further parameters passed to the function mtext (only if 'add.text = TRUE').
arg.points	a named list of graphical parameters passed to the function points .
...	additional graphical arguments passed to the functions plot , contour or filled.contour .

Details

plot.GoF is the plotting method function of an R object of class 'GoF', that is, the output of a goodness-of-fit function (see [AIC.cglasso](#), or [BIC.cglasso](#)). This function produces a plot aimed both to evaluate the sequence of fitted models in terms of goodness-of-fit and to identify the optimal values of the tuning parameters.

If a tuning parameter is held fixed, then plot.GoF produces a plot showing the chosen measure of goodness-of-fit as a function of the remaining tuning parameter. In this case, the optimal value is identified by a vertical dashed line. The degrees-of-freedom of the selected fitted model are also shown.

If the cglasso model is fitted using both a sequence of ρ and λ values, then plot.GoF produces a contour plot and a triangle is used to identify the optimal pair of the two tuning parameters.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[cglasso](#), [AIC.cglasso](#), [BIC.cglasso](#), [summary.cglasso](#) and [select.cglasso](#).

Examples

```
set.seed(123)
n <- 1000L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z, nlambda = 1L)
plot(AIC(out))
plot(BIC(out))

out <- cglasso(. ~ ., data = Z, nrho = 1L)
plot(AIC(out))
plot(BIC(out))

out <- cglasso(. ~ ., data = Z)
plot(AIC(out))
plot(BIC(out))
```

predict

Predict Method for cglasso and cggm Fits

Description

Obtains predictions from an R object inheriting class ‘cglasso’.

Usage

```
## S3 method for class 'cglasso'
predict(object, type = c("B", "mu", "Sigma", "Theta"), X.new, lambda.new, rho.new,
        ...)

## S3 method for class 'cggm'
predict(object, X.new, ...)
```

Arguments

object an R object inheriting class ‘cglasso’, that is, the output of the model-fitting function ‘cglasso’ or ‘cggm’.

type	a description of prediction required.
X.new	matrix of new values for X at which predictions are to be made. This argument is used only if 'type = "mu"'.
lambda.new	value of the tuning parameter λ at which predictions are required.
rho.new	value of the tuning parameter ρ at which predictions are required.
...	further arguments passed to or from other methods.

Details

If object has S3 class 'cglasso', then for a new pair of the tuning parameters λ and ρ , the predict function can be used to predict the estimate of the regression coefficient matrix ('type = "B"'), the estimate of the covariance matrix ('type = "Sigma"') or the estimate of the precision matrix ('type = "Theta"'). If X.new is missing and 'type = "mu"', then the predict function returns the predicted values using the matrix of predictors X, otherwise the predicted fitted values are computed using the matrix X.new.

For a new pair of the tuning parameters λ and ρ , the predicted values are computed using a bilinear interpolation.

If the object has S3 class 'cggm', then the predict function returns only the predicted fitted values using the argument X.new.

Value

The matrix of predicted values.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

Model-fitting function [cglasso](#) and the other accessor functions [coef.cglasso](#), [fitted.cglasso](#), [residuals.cglasso](#) and [impute](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z)
rho.new <- mean(out$rho)
Theta.pred <- predict(out, rho.new = rho.new, type = "Theta")
Theta.pred

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
```

```

n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z)
rho.new <- mean(out$rho)
lambda.new <- mean(out$lambda)
Theta.pred <- predict(out, lambda.new = lambda.new, rho.new = rho.new, type = "Theta")
Theta.pred

```

QFun

*Extract Q-Function***Description**

‘QFun’ extracts the values of the Q-function from an R object inheriting class ‘cglasso’.

Usage

```
QFun(object, mle, verbose = FALSE, ...)
```

Arguments

object	an R object inheriting class ‘cglasso’, that is, the output of the model-fitting functions cglasso and cggm .
mle	logical. Should Q-values be computed using the maximum likelihood estimates? Default depends on the class of the argument object: <code>mle = FALSE</code> for objects with class <code>cglasso</code> and <code>mle = TRUE</code> for objects with class <code>cggm</code> .
verbose	logical for printing out a progress bar on the R console. Default is <code>verbose = FALSE</code> .
...	further arguments passed to cggm .

Details

‘QFun’ returns the value of the Q-function, i.e., the value of the function maximised in the M-step of the EM algorithm. The Q-function is defined as follows:

$$\frac{n}{2} \{ \log \det \Theta - \text{tr}(S\Theta) - p \log(2\pi) \},$$

where S is the ‘working’ empirical covariance matrix computed during the E-step.

QFun is used as a workhorse function to compute the measures of goodness-of-fit returned by the functions [AIC.cglasso](#) and [BIC.cglasso](#).

The function ‘`print.QFun`’ is used to improve the readability of the results.

Value

'QFun' returns an R object of S3 class "QFun", i.e., a named list containing the following components:

value	a matrix with the values of the Q-function.
df	a matrix with the number of estimated non-zero parameters.
dfB	a matrix with the number of estimated non-zero regression coefficients.
dfTht	a matrix with the number of estimated non-zero partial correlation coefficients.
n	the sample size.
p	the number of response variables.
q	the number of columns of the design matrix X used to fit the model.
lambda	the λ -values used to fit the model.
nlambda	the number of λ -values.
rho	the ρ -values used to fit the model.
nrho	the number of ρ -values.
model	a description of the fitted model passed through the argument object.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[AIC.cglasso](#), [BIC.cglasso](#), [cglasso](#), [cggm](#), [summary.cglasso](#), [select.cglasso](#) and [to_graph](#).

Examples

```
set.seed(123)
# Y ~ N(b0+ XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1:p, 1:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.5,
          probna = 0.05)
out <- cglasso(. ~ ., data = Z)
QFun(out)

out.mle <- cggm(out, lambda.id = 3L, rho.id = 3L)
QFun(out.mle)
```

qqcnorm *Quantile-Quantile Plots for a datacggm Object*

Description

Creates a quantile-quantile plot for a censored Gaussian distribution.

Usage

```
qqcnorm(x, which, max.plot = 1L, save.plot = FALSE, grdev = pdf, grdev.arg,
        main = "Censored Normal Q-Q Plot", xlab = "Theoretical Quantiles",
        ylab = "Sample Quantiles", plot.it = TRUE, plot.pch = c(2L, 20L),
        plot.col = c(2L, 1L), plot.cex = c(2L, 1L), abline = FALSE,
        line.col = "gray50", line.lwd = 1L, line.lty = 2L, ...)
```

Arguments

x	an object of class 'datacggm'.
which	a vector of integers used to specify the response variables for which the histogram is required.
max.plot	the maximum number of plots drawn in a single figure.
save.plot	a logical variable or a string specifying the path of the directory where plots will be saved. Letting 'save.plot = TRUE', the required plots will be saved as external files inside the current working directory. User can save these files on a specific working directory passing the absolute path through the argument save.plot. On exit, working directory will be always set the previous one.
grdev	the graphics device used to save the required histograms on external files. See 'device' for more details.
grdev.arg	additional parameters passed to the graphics device specified by 'grdev'.
main, xlab, ylab	plot labels.
plot.it	logical. Should the result be plotted?
plot.pch	a pair of graphical parameters. The first entry specifies the symbol used for plotting the points associated to the censoring values <i>lo</i> and <i>up</i> (by default a triangle), whereas the second entry specifies the symbol used for the remaining points (by default a black point).
plot.col	a pair of graphical parameters. The first entry specifies the colour used for plotting the points associated to the censoring values <i>lo</i> and <i>up</i> (by default a red triangle), whereas the second entry specifies the colour used for the remaining points (by default a black point).
plot.cex	a pair of graphical parameters. The first entry specifies the size of the symbol used for plotting the points associated to the censoring values <i>lo</i> and <i>up</i> , whereas the second entry specifies the size of the symbol used for the remaining points.
abline	logical. Should the line $y = x$ be plotted?

<code>line.col</code>	graphical parameter. If <code>'abline = TRUE'</code> , then this argument specifies the colour of the line $y = x$.
<code>line.lwd</code>	graphical parameter. If <code>'abline = TRUE'</code> , then this argument specifies the line width of the line $y = x$.
<code>line.lty</code>	graphical parameter. If <code>'abline = TRUE'</code> , then this argument specifies the line type of the line $y = x$.
<code>...</code>	additional graphical parameter passed to <code>'plot'</code> .

Details

`'qqcnorm'` produces a censored normal QQ plot, that is, a graphical method for comparing the empirical distribution of a given response variable (specified by the argument `which`) to the censored Gaussian distribution, which is defined as:

$$\begin{aligned} \Phi((lo - \mu)/\sigma) & \quad \text{if } y \leq lo \\ \phi((y - \mu)/\sigma)/\sigma & \quad \text{if } lo < y < up \\ 1 - \Phi((up - \mu)/\sigma) & \quad \text{if } y \geq up \end{aligned}$$

where ϕ and Φ are the probability density function and the cumulative distribution of the standard normal distribution, respectively, whereas lo and up are the lower and upper censoring values, respectively.

The comparison is done by plotting the empirical quantiles (y -coordinate) against the theoretical quantiles (x -coordinate) of the censored Gaussian distribution, which are defined as follows:

$$\begin{aligned} lo & \quad \text{if } p \leq \Phi((lo - \mu)/\sigma) \\ \mu + \sigma\Phi^{-1}(p) & \quad \text{if } \Phi((lo - \mu)/\sigma) < p < 1 - \Phi((up - \mu)/\sigma) \\ up & \quad \text{if } p \geq 1 - \Phi((up - \mu)/\sigma) \end{aligned}$$

where $p \in (0, 1)$. If the two distributions are similar, the points will approximately lie on the line $y = x$. If the distributions are linearly related, the points will approximately lie on a line, but not necessarily on the line $y = x$. In order to evaluate if the proportions of left/right-censored values are similar to the Gaussian tail probabilities, points corresponding to the censored values are plotted using a specific symbol (see argument `'plot.pch'`), colour (see argument `'plot.col'`) and size (see argument `'plot.cex'`).

Finally, maximum likelihood estimates of the marginal parameters μ and σ are computed as described in `'datacggm'` and can be extracted from an R of class `'datacggm'` by using the functions `'ColMeans'` and `'ColVars'`, respectively.

Value

A named list is silently returned. Each element of the list contains a two-columns matrix; first columns (named `'x'`) contains the theoretical quantiles whereas second columns (named `'y'`) contains the empirical quantiles.

Author(s)

Gianluca Sottile (<gianluca.sottile@unipa.it>)

References

Augugliaro, L., Sottile, G., and Vinciotti, V. (2020) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.

Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#), [rcggm](#), [ColMeans](#), [ColVars](#) and [hist.datacggm](#).

Examples

```
set.seed(123)

# a dataset from a right-censored Gaussian graphical model
n <- 1000L
p <- 10L
Y <- matrix(rnorm(n * p), n, p)
up <- 1
Y[Y >= up] <- up
Z <- datacggm(Y = Y, up = up)
qqcnorm(Z, max.plot = 4L)

# a dataset from a conditional censored Gaussian graphical model
n <- 1000L
p <- 10L
q <- 2
Y <- matrix(rnorm(n * p), n, p)
up <- 1
lo <- -1
Y[Y >= up] <- up
Y[Y <= lo] <- lo
X <- matrix(rnorm(n * q), n, q)
Z <- datacggm(Y = Y, lo = lo, up = up, X = X)
qqcnorm(Z, max.plot = 4L)
```

rcggm

Simulate Data from a Conditional Gaussian Graphical Model with Censored and/or Missing Values

Description

‘rcggm’ function is used to produce one or more samples from a conditional Gaussian graphical model with censored and/or missing values.

Usage

```
rcggm(n, p, b0, X, B, Sigma, probl, probr, probna, ...)
```

Arguments

n	the number of samples required (optional, see below for a description).
p	the number of response variables (optional, see below for a description).
b0	a vector of length p used to specify the intercepts. Default is a zero vector of length p (optional, see below for a description).
X	a matrix of dimension $n \times q$ used to model the expected values of the response variables (optional, see below for a description).
B	a matrix of dimension $q \times p$ used to specify the regression coefficients. If X is missing then B is set equal to NULL (optional, see below for a description).
Sigma	a positive-definite symmetric matrix specifying the covariance matrix of the response variables. Default is the identity matrix (optional, see below for a description).
probl	a vector giving the probabilities that the response variables are left-censored.
probr	a vector giving the probabilities that the response variables are right-censored.
probna	the probability that a response value is missing-at-random. By default 'probna' is set equal to zero.
...	further arguments passed to the function 'mvrnorm'.

Details

'The rcggm' function simulates a dataset from a conditional Gaussian graphical model with censored or missing values and returns an object of class 'datacggm'. Censoring values are implicitly specified using arguments probl and probr, that is, lo and up are computed in such a way that the average probabilities of left and right censoring are equal to probl and probr, respectively. Missing-at-random values are simulated using a Bernoulli distribution with probability probna.

The dataset is simulated through the following steps:

1. lower and upper censoring values (lo and up) are computed according to the arguments probl and probr;
2. The function `mvrnorm` is used to simulate one or more samples from the multivariate Gaussian distribution specified by the arguments b0, X, B and Sigma;
3. The response values that are outside of the interval [lo, up] are replaced with the corresponding censoring values;
4. if probna is greater than zero, then missing-at-random values are simulated using a Bernoulli distribution with probability probna.

Model	n	p	b0	X	B	Sigma	Gaussian distribution
1	x	x					$Y \sim N(0, I)$
2	x					x	$Y \sim N(0, \Sigma)$
3	x		x				$Y \sim N(b0, I)$
4	x		x			x	$Y \sim N(b0, \Sigma)$
5				x	x		$Y \sim N(XB, I)$
6				x	x	x	$Y \sim N(XB, \Sigma)$
7			x	x	x		$Y \sim N(b0 + XB, I)$
8			x	x	x	x	$Y \sim N(b0 + XB, \Sigma)$

The previous table sums up the default setting of the multivariate Gaussian distribution used in step 2 (specified arguments are marked by the symbol ‘x’). See also below for some examples.

Value

rcggm returns an object of class ‘datacggm’. See [datacggm](#) for further details.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

Augugliaro, L., Sottile, G., and Vinciotti, V. (2020a) <[doi:10.1007/s11222020099457](https://doi.org/10.1007/s11222020099457)>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.

Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020b) <[doi:10.1093/biostatistics/kxy043](https://doi.org/10.1093/biostatistics/kxy043)>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#).

Examples

```
set.seed(123)

n <- 100
p <- 3
q <- 2
b0 <- rep(1, p)
X <- matrix(rnorm(n * q), n, q)
B <- matrix(rnorm(q * p), q, p)
Sigma <- outer(1:p, 1:p, function(i, j) 0.3^abs(i - j))
probl <- 0.05
probr <- 0.05
probna <- 0.05

# Model 1: Y ~ N(0, I)
Z <- rcggm(n = n, p = p, probl = probl, probr = probr, probna = probna)
summary(Z)

# Model 2: Y ~ N(0, Sigma)
Z <- rcggm(n = n, Sigma = Sigma, probl = probl, probr = probr, probna = probna)
summary(Z)

# Model 3: Y ~ N(b0, I)
Z <- rcggm(n = n, b0 = b0, probl = probl, probr = probr, probna = probna)
summary(Z)

# Model 4: Y ~ N(b0, Sigma)
Z <- rcggm(n = n, b0 = b0, Sigma = Sigma, probl = probl, probr = probr,
           probna = probna)
```

```
summary(Z)

# Model 5:  $Y \sim N(XB, I)$ 
Z <- rcggm(X = X, B = B, probl = probl, probr = probr, probna = probna)
summary(Z)

# Model 6:  $Y \sim N(XB, \text{Sigma})$ 
Z <- rcggm(X = X, B = B, Sigma = Sigma, probl = probl, probr = probr,
           probna = probna)
summary(Z)

# Model 7:  $Y \sim N(b_0 + XB, I)$ 
Z <- rcggm(b0 = b0, X = X, B = B, probl = probl, probr = probr, probna = probna)
summary(Z)

# Model 8:  $Y \sim N(b_0 + XB, \text{Sigma})$ 
Z <- rcggm(b0 = b0, X = X, B = B, Sigma = Sigma, probl = probl, probr = probr,
           probna = probna)
summary(Z)
```

residuals

Extract Model Residuals

Description

Extracts model residuals from an R object inheriting class 'cglasso'.

Usage

```
## S3 method for class 'cglasso'
residuals(object, type = c("observed", "working"), lambda.id, rho.id,
          drop = TRUE, ...)
```

Arguments

object	an R object inheriting class 'cglasso', that is, the output of the model-fitting functions cglasso and cggm .
type	a description of the desired residuals (see section 'Details').
lambda.id	a vector of integers used to specify the λ -values.
rho.id	a vector of integers used to specify the ρ -values.
drop	logical. Dimensions can only be dropped if their extent is one.
...	further arguments passed to or from other methods.

Details

The accessor function ‘residuals’ returns an array storing the ‘observed’ or ‘working’ residuals, depending on the argument type.

The ‘observed’ residuals are defined as the difference between the observed response values and the corresponding fitted expected values. For missing and censored response values, the ‘observed’ residuals are set equal to ‘NA’.

The ‘working’ residuals are obtained as a byproduct of the EM-algorithm used to fit the model. In the E-step, the algorithm computes the ‘working’ response matrix, that is, the response matrix with missing values replaced by the conditional expected values of the multivariate normal distribution and censored values replaced by the conditional expected values of the multivariate truncated normal distribution. The ‘working’ residuals are defined as the difference between ‘working’ responses and fitted expected values.

Value

The accessor function ‘residuals’ returns an array storing the desired residuals.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

Model-fitting functions [cglasso](#), [cggm](#) and the accessor functions [coef.cglasso](#), [fitted.cglasso](#), [predict.cglasso](#) and [impute](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
residuals(out, type = "observed", rho.id = 3L, drop = TRUE)
residuals(out, type = "working", rho.id = 3L, drop = TRUE)

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 1000L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
```

```
residuals(out, type = "observed", lambda.id = 3L, rho.id = 3L, drop = TRUE)
residuals(out, type = "working", lambda.id = 3L, rho.id = 3L, drop = TRUE)
```

rowNames + colNames *Row and Column Names of a "datacggm" Object*

Description

Retrieve or set the row or column names of a "datacggm" object.

Usage

```
rowNames(x)
rowNames(x) <- value

colNames(x)
colNames(x) <- value
```

Arguments

x an R object of class 'datacggm'.
value a named list with elements 'X' and 'Y'.

Details

For an R object of class 'datacggm', rowNames (colNames) retrieves or set rownames (colnames) for matrices Y and X. See below for some examples.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[datacggm](#), [rcggm](#) and the method function [dimnames](#).

Examples

```
set.seed(123)
# a dataset from a censored Gaussian graphical model
n <- 100L
p <- 3L
Z <- rcggm(n = n, p = p, probl = 0.05, probr = 0.05)

rowNames(Z)
rowNames(Z) <- list(Y = paste0("i", seq_len(n)))
# the same as rowNames(Z)$Y <- paste0("i", seq_len(n))
rowNames(Z)
```

```

colNames(Z)
colNames(Z) <- list(Y = paste("Y", 1:p, sep = ":"))
# the same as colNames(Z)$Y <- paste("Y", 1:p, sep = ":")
colNames(Z)

# a dataset from a conditional censored Gaussian graphical model
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
Z <- rcggm(n = n, b0 = b0, X = X, B = B, probl = 0.05, probr = 0.05)

rowNames(Z)$Y <- paste0("i", seq_len(n))
rowNames(Z)$X <- paste0("i", seq_len(n))
dimnames(Z)

colNames(Z)$Y <- paste("Y", 1:p, sep = ":")
colNames(Z)$X <- paste("X", 1:q, sep = ":")
dimnames(Z)

```

select.cglasso

Model Selection for the Conditional Graphical Lasso Estimator

Description

‘select.cglasso’ returns the optimal fitted model selected by a chosen measure of goodness-of-fit.

Usage

```
select.cglasso(object, GoF = AIC, ...)
```

Arguments

object	an R object of class cglasso.
GoF	a valid goodness-of-fit function, such as AIC.cglasso or BIC.cglasso , or an R object of class ‘GoF’.
...	further arguments passed to the chosen goodness-of-fit function (argument ‘GoF’).

Details

The function `select.cglasso` evaluates the goodness-of-fit of the models fitted by `cglasso` and extracts the selected model.

Model evaluation can be made in two ways. The easiest way is to use a valid goodness-of-fit function, such as [AIC.cglasso](#) or [BIC.cglasso](#). In this case, further arguments are passed to these functions by ‘...’. The second way consists on passing the output of a goodness-of-fit function,

that is, an R object of class ‘GoF’. Usually, this approach is preferable when the computation of the chosen goodness-of-fit measure is time-consuming, such as when the sample size is small relative to the number of parameters and the AIC or BIC functions are used to evaluate a long sequence of fitted models. In these cases, we suggest the computation of several measures of goodness-of-fit in a preliminary step and then the use of the `select.cglasso` function in a subsequent step to select the optimal fitted model.

Value

‘`select.cglasso`’ returns the optimal fitted model.

Author(s)

Gianluca Sottile (<gianluca.sottile@unipa.it>)

See Also

[cglasso](#), [AIC.cglasso](#) and [BIC.cglasso](#).

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z)
select.cglasso(out) # models selection by AIC
select.cglasso(out, GoF = BIC) # models selection by BIC
select.cglasso(out, GoF = BIC, mle = TRUE, g = 0.5) # models selection by eBIC

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z, lambda = 0.01)
select.cglasso(out) # models selection by AIC
select.cglasso(out, GoF = BIC) # models selection by BIC
select.cglasso(out, GoF = BIC, mle = TRUE, g = 0.5) # models selection by eBIC

out <- cglasso(. ~ ., data = Z, rho = 0.01)
select.cglasso(out) # models selection by AIC
```

```

select.cglasso(out, GoF = BIC)           # models selection by BIC
select.cglasso(out, GoF = BIC, mle = TRUE, g = 0.5) # models selection by eBIC

out <- cglasso(. ~ ., data = Z)
select.cglasso(out)                     # models selection by AIC
select.cglasso(out, GoF = BIC)         # models selection by BIC
select.cglasso(out, GoF = BIC, mle = TRUE, g = 0.5) # models selection by eBIC

```

ShowStructure *Show Package Structure*

Description

ShowStructure gives information about the package structure.

Usage

```
ShowStructure(module = c("ALL", "DM", "MF", "MS", "NA"), description = TRUE,
              plot = TRUE)
```

Arguments

module	a description of the module for which the description is required.
plot	logical. If TRUE a graph showing the package structure is plotted.
description	logical. If TRUE a description of the available function is printed on screen.

Value

ShowStructure silently returns a named list with components:

description	a named list containing the description of the available functions.
graph	an R object of class 'igraph' describing the package structure.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

summary.cglasso

Summarizing cglasso and cggm Fits

Description

'summary' produces a summary of the models fitted using cglasso or cggm.

Usage

```
## S3 method for class 'cglasso'
summary(object, GoF = AIC, print.all = TRUE, digits = 3L, ...)
```

Arguments

object	an R object inheriting class cglasso, that is, the output of the model-fitting functions cglasso and cggm .
GoF	a valid goodness-of-fit function, such as AIC.cglasso or BIC.cglasso , or an R object of class 'GoF'.
print.all	logical. Should all summary statistics be printed?
digits	the minimum number of significant digits to be used: see ' print.default '.
...	the <i>penalty</i> parameter passed to the goodness-of-fit function (argument 'GoF') or further arguments passed to ' print.data.frame ' and ' print.listof '.

Details

The function `summary.cglasso` computes the summary statistics needed to evaluate the goodness-of-fit of the models fitted by `cglasso` or `cggm`.

Model evaluation can be made in two ways. The easiest way is to use a valid goodness-of-fit function, such as [AIC.cglasso](#) or [BIC.cglasso](#). In this case, further arguments can be passed to these functions by the argument '...'. The second way consists in passing the output of a goodness-of-fit function, that is, an R object of class 'GoF'. Usually, this approach is preferable when the computation of the chosen goodness-of-fit measure is time-consuming, such as when the sample size is small relative to the number of parameters and the AIC or BIC functions are used to evaluate a sequence of fitted models. In these cases, we suggest the computation of several measures of goodness-of-fit in a preliminary step and then the use of `summary.cglasso` in a subsequent step to evaluate the sequence of fitted models.

To improve the readability of the results, the output is divided into two sections.

The first section is structured in a sequence of tables showing, for each combination of the two tuning parameters 'lambda' and 'rho', the number of estimated non-zero regression coefficients ('df.B'), the number of estimated non-zero partial correlation coefficients ('df.Tht'), the degrees-of-freedom ('df') and the value of the goodness-of-fit measure used to evaluate the fitted models. To help the user with the identification of the optimal fitted model, the last column of each table reports the ranking of the models (where the optimal model is marked with the symbol '<-').

The second section reports the summary statistics of the selected optimal model.

Value

A named list with the following elements is silently returned:

table	data.frame containing the summary statistics used to evaluate the sequence of fitted models.
lambda.id	position of the optimal λ -value identified by the chosen goodness-of-fit function.
rho.id	position of the optimal ρ -value identified by the chosen goodness-of-fit function.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

See Also

[cglasso](#), [cggm](#), [AIC.cglasso](#) and [BIC.cglasso](#).

Examples

```
set.seed(123)

# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z)
summary(out) # models evaluation by AIC
summary(out, GoF = BIC) # models evaluation by BIC
summary(out, GoF = BIC, mle = TRUE, g = 0.5) # models evaluation by eBIC

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)

out <- cglasso(. ~ ., data = Z, lambda = 0.01)
summary(out) # models evaluation by AIC
summary(out, GoF = BIC) # models evaluation by BIC
summary(out, GoF = BIC, mle = TRUE, g = 0.5) # models evaluation by eBIC

out <- cglasso(. ~ ., data = Z, rho = 0.01)
summary(out) # models evaluation by AIC
summary(out, GoF = BIC) # models evaluation by BIC
```

```
summary(out, GoF = BIC, mle = TRUE, g = 0.5) # models evaluation by eBIC

out <- cglasso(. ~ ., data = Z)
summary(out) # models evaluation by AIC
summary(out, GoF = BIC) # models evaluation by BIC
summary(out, GoF = BIC, mle = TRUE, g = 0.5) # models evaluation by eBIC
```

summary.datacggm *Summarizing Objects of Class 'datacggm'*

Description

The function 'summary.datacggm' produces summaries of an object of class 'datacggm'.

Usage

```
## S3 method for class 'datacggm'
summary(object, n, quantile.type = 7L, digits = 3L, quote = FALSE, ...)
```

Arguments

object	an object of class 'datacggm' for which a summary is desired.
n	maximum number of rows printed on video. By default all rows are printed.
quantile.type	an integer between 1 and 9 selecting one of the nine quantile algorithms (see also quantile function).
digits	integer, used for number formatting with 'format()'. By default, digits are 3.
quote	logical, indicating whether or not strings should be printed with surrounding quotes.
...	further arguments passed to print.listof .

Details

The function 'summary.datacggm' extends the results given by 'summary.matrix()' taking into account the specific structure of an object of class 'datacggm'. Summary statistics are printed out in two sections: the first section reports the summary statistics for the response matrix 'Y', whereas the second section is devoted to summarising the matrix of predictors 'X'.

For each response variable, the first section reports the standard summary statistics computed using only the observed values, that is the entries of the response matrix whose status indicators are equals to zero (see 'event' for more details), together with the information about the lower and upper censoring values (columns named Lower and Upper) and the percentage of missing-at-random, left-censored and right-censored response values (columns named NA%, LC% and RC%, respectively).

The second section is divided into two subsections reporting the main summary statistics computed for numeric and categorical predictors, respectively.

The two sections are printed out in such a way that the readability of the summary statistics is improved in a high-dimensional setting, that is, when the number of predictors and response variables exceeds the sample size.

Value

'summary.datacggm' returns a named list with components

- Y a matrix with class 'table' obtained by computing the summary measures on the response variables.
- X.numeric a matrix with class 'table' obtained by computing the summary measures on the numeric predictors.
- X.categorical a list containing the summary measures on the categorical predictors.

Author(s)

Luigi Augugliaro (<luigi.augugliaro@unipa.it>)

References

- Augugliaro, L., Sottile, G., and Vinciotti, V. (2020a) <doi:10.1007/s11222020099457>. The conditional censored graphical lasso estimator. *Statistics and Computing* **30**, 1273–1289.
- Augugliaro, L., Abbruzzo, A., and Vinciotti, V. (2020b) <doi:10.1093/biostatistics/kxy043>. ℓ_1 -Penalized censored Gaussian graphical model. *Biostatistics* **21**, e1–e16.

See Also

[datacggm](#), [rcggm](#) and [event](#).

Examples

```
set.seed(123)

# case 1: Y ~ N(0, Sigma)
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random equals to 0.05
n <- 50L
p <- 100L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05, probna = 0.05)
summary(Z, max = n * p)

# case 2: Y ~ N(b0 + XB, Sigma) and
# 1. probability of left/right censored values equal to 0.05
# 2. probability of missing-at-random equals to 0.05
n <- 50L
p <- 100L
q <- 100L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05,
```

```

summary(Z)
summary(Z, max = n * p)

```

to_graph

Create Graphs from cglasso or cggm Objects

Description

'to_graph' returns a named list of graphs using the results of an R object of class 'cglasso' or 'cggm'.

Usage

```

to_graph(object, GoF = AIC, lambda.id, rho.id, weighted = FALSE, simplify = TRUE,
        ...)

```

Arguments

object	an R object inheriting class 'cglasso', that is, the output of the model-fitting functions cglasso and cggm .
GoF	a valid goodness-of-fit function, such as AIC.cglasso or BIC.cglasso , or an R object of class 'GoF'.
lambda.id	optional. If object has class 'cglasso', this argument is an integer used to identify a specific fitted cglasso model, otherwise (i.e. if the object has class 'cggm') this argument can be omitted. See section 'Details' for more details.
rho.id	optional. If object has class 'cglasso', this argument is an integer used to identify a specific fitted cglasso model, otherwise (i.e. if object has class 'cggm') this argument can be omitted. See section 'Details' for more details.
weighted	logical. Should weighted graphs be created? Default is FALSE.
simplify	logical. Should isolated vertices be removed from the graph? Default is TRUE, i.e., isolated vertices are removed.
...	further arguments passed to the chosen goodness-of-fit function (argument 'GoF').

Details

'to_graph' returns a named list of graphs using the results of an R object of class 'cglasso' or 'cggm'.

If object has class 'cglasso', then the goodness-of-fit function passed through the argument GoF is used to identify the adjacency matrix (`object$InfoStructure$Adj_yy`) describing the undirected edges among the p response variables. If the model is fitted using q predictors, then the matrix describing the effects of the predictors onto the response variables (see `object$InfoStructure$Adj_xy`) is also returned. Finally, these matrices are used to return an undirected and directed graph. Optionally, the user can identify a specific fitted model using the arguments `lambda.id` and `rho.id`.

If object has class 'cggm', then `GoF`, `lambda.id` and `rho.id` can be omitted.

If argument `weighted` is set equal to `'TRUE'`, then the estimated precision matrix and, if available, the estimated regression coefficient matrix are used to return weighted graphs. In this case, edges associated with positive estimates are shown using a solid line. Otherwise, a dashed line is used.

Value

`'to_graph'` returns an R object of S3 class `"cglasso2igraph"`, i.e., a named list containing the following components:

<code>Gyy</code>	an undirected graph representing the conditional dependence structure among the p response variables.
<code>Gxy</code>	a directed graph representing the effects of the q predictors onto the p response variables.

Each component is an R object of class `igraph`.

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See Also

`cglasso`, `cggm` and `plot.cglasso2igraph`. For more details about the object of class `'igraph'`, the interested reader is referred to the package `igraph`.

Examples

```
set.seed(123)
# Y ~ N(0, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out)
out.graph

# Y ~ N(b0 + XB, Sigma) and probability of left/right censored values equal to 0.05
n <- 100L
p <- 3L
q <- 2L
b0 <- runif(p)
B <- matrix(runif(q * p), nrow = q, ncol = p)
X <- matrix(rnorm(n * q), nrow = n, ncol = q)
rho <- 0.3
Sigma <- outer(1L:p, 1L:p, function(i, j) rho^abs(i - j))
Z <- rcggm(n = n, b0 = b0, X = X, B = B, Sigma = Sigma, probl = 0.05, probr = 0.05)
out <- cglasso(. ~ ., data = Z)
out.graph <- to_graph(out, lambda.id = 3, rho.id = 3, weighted = TRUE)
out.graph
```

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