Package ‘sglasso’

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Description RCON(V, E) models are a kind of restriction of the Gaussian Graphical Models defined by a set of equality constraints on the entries of the concentration matrix. 'sglasso' package implements the structured graphical lasso (sglasso) estimator proposed in Abbruzzo et al. (2014) for the weighted 11-penalized RCON(V, E) model. Two cyclic coordinate algorithms are implemented to compute the sglasso estimator, i.e. a cyclic coordinate minimization (CCM) and a cyclic coordinate descent (CCD) algorithm.
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**Description**

*RCON*(V, E) models (Hojsgaard, et al., 2008) are a kind of restriction of the Gaussian Graphical Models defined by a set of equality constraints on the entries of the concentration matrix. *sglasso* package implements the structured graphical lasso (*sglasso*) estimator proposed in Abbruzzo et al. (2014) for the weighted 11-penalized *RCON*(V, E) model. Two cyclic coordinate algorithms are implemented to compute the *sglasso* estimator, i.e. a cyclic coordinate minimization (CCM) and a cyclic coordinate descent (CCD) algorithm.

**Details**

- **Package:** *sglasso*
- **Type:** Package
- **Version:** 1.2.3
- **Date:** 2018-05-26
- **License:** GPL (>=2)

**Author(s)**

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**References**


L1-penalized Factorial Graphical Lasso Model

Description

Fit the weight l1-penalized factorial dynamic Gaussian Graphical Model.

Usage

`fglasso(s, model, tp, p, ...)`

Arguments

- `s`: the empirical variance/covariance matrix;
- `model`: a list or a matrix used to specify the factorial dynamic Gaussian Graphical Model (see Details);
- `tp`: number of time points;
- `p`: number of random variables observed for each time point;
- `...`: further arguments passed to `sglasso`.

Details

The factorial dynamic Gaussian Graphical Model (Abbruzzo et al., 2015) is a special kind of RCON(V, E) model (Hojsgaard, et al., 2008) proposed to study dynamic networks. Let $X_t = (X_{it},\ldots,X_{it})'$ be a p-dimensional random variable at time $t$. Assuming that $X = (X_1',\ldots,X_T')$ follows a multivariate normal distribution, the concentration matrix $K$ has the following block structure

$$K = \begin{pmatrix}
K_{1,1} & K_{1,2} & \ldots & K_{1,T} \\
K_{2,1} & K_{2,2} & \ldots & K_{2,T} \\
\vdots & \vdots & \ddots & \vdots \\
K_{T,1} & K_{T,2} & \ldots & K_{T,T}
\end{pmatrix},$$

where $K_{t,t}$ give information about the conditional independence structure among the p random variables at time $t$, and $K_{t,t+h}$ give information about the conditional independence structure between $X_t$ and $X_{t+h}$. An interpretation of the elements of the submatrices $K_{t,t+h}$ brings to the notion of natural structure, i.e.,

$$K_{t,t+h} = \begin{pmatrix}
k_{1,1}^{t,t+h} & 0 & \ldots & 0 \\
0 & k_{2,2}^{t,t+h} & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \ldots & k_{p,p}^{t,t+h}
\end{pmatrix} + \begin{pmatrix}
0 & k_{1,2}^{t,t+h} & \ldots & k_{1,p}^{t,t+h} \\
k_{2,1}^{t,t+h} & 0 & \ldots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
k_{p,1}^{t,t+h} & k_{p,2}^{t,t+h} & \ldots & 0
\end{pmatrix}.$$

The entries of the first matrix are called self-self conditional dependences at temporal lag $h$ and represent the (negative) self-similarity of a given random variable across different time points. The entries of the second matrix are the conditional dependence among the p random variables. To make the interpretation of the results more relevant and, at the same time, reduce the number of parameters, the authors propose the following equality constraints:
Argument model is used to specify the restrictions previously described. This argument can be a named list or a matrix with dimension \( nlag \times 2 \), where \( nlag \leq tp \). To gain more insight, suppose that we want to model only the sub-matrices \( K_{t,t} \) and \( K_{t,t+1} \), i.e., the sub-matrices corresponding to the temporal lag zero and one. A possible R code is

```r
model.m <- matrix("", nrow = 2, ncol = 2)
rownames(model.m) <- c("lag0", "lag1")
colnames(model.m) <- c("s", "n")
model.m[1, ] <- c("c", "ut")
model.m[2, ] <- c("t", ".")
```

In this example we are modelling the diagonal elements of the sub-matrices \( K_{t,t} \) with the constant effect while the off-diagonal elements are modelled by the interaction effect. In the same way, the diagonal elements of the sub-matrices \( K_{t,t+1} \) are modelled by the time effect while the remaining elements are equal to zero. The fglasso function passes the matrix `model.m` to the internal function `fglasso_modelRmask`, i.e.,

```r
mask <- fglasso_modelRmask(model.m, tp)
```

which returns the mask used in sglasso to fit the specified factorial dynamic Gaussian Graphical model. The same model can be specified by the following named list

```r
model.list <- list(lag0 = c(s = "c", n = "ut"), lag1 = c(s = "t", n = "."))
```

See the example below for more details.

**Value**

fglasso returns an object with S3 class "sglasso". See the corresponding manual for more details.

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**References**


See Also

sgllasso function.

Examples

```
# fglasso solution path
N <- 50
tp <- 3
p <- 3
X <- matrix(rnorm(N * p * tp), N, tp * p)
S <- crossprod(X) / N
model <- list(lag0 = c(s = "c", n = "ut"), lag1 = c(s = "t", n = "."))
out.fglasso <- fglasso(S = S, model = model, tp = tp, p = p)
out.fglasso
```

---

**gplot**

*Plotting Sparse Graph*

**Description**

gplot is a generic function for plotting sparse graphs.

**Usage**

gplot(object, ...)

**Arguments**

- **object**: fitted sgllasso/fglasso object;
- **...**: other parameters passed to `gplot.sglasso` or `gplot.fglasso`.

**Details**

gplot is a generic function used to plot a graph estimated by sgllasso or fglasso. See the method function `gplot.sglasso` or `gplot.fglasso` for more details about the specific arguments.
**gplot.fglasso**

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**See Also**

gplot.sglasso and gplot.fglasso method functions.

---

**gplot.fglasso**

*Plotting Sparse Factorial Dynamic Gaussian Graphical Model*

**Description**

gplot.fglasso shows the sequence of graphs estimated by fglasso.

**Usage**

```r
## S3 method for class 'fglasso'
gplot(object, rhoid, tp = c(1, 2), sub.tp1, sub.tp2, cex.sub = 1,
      k = 1.5, layout = layout.circle, ...)
```

**Arguments**

- `object`: fitted fglasso object;
- `rhoid`: an integer used to specify the \( \rho \)-value used to fit the fglasso model;
- `tp`: a vector of length equal to two used to specify the time points of the two graphs that will be compared. By default the first two time points are used;
- `sub.tp1`: sub title for the graph estimated at time point `tp[1]`;
- `sub.tp2`: sub title for the graph estimated at time point `tp[2]`;
- `cex.sub`: a numerical value giving the amount by which plotting sub titles should be magnified relative to the default;
- `k`: value used to specify the distance between the two graphs;
- `layout`: a function or a matrix used to specify the layout of the graphs that will be plotted. By default the `layout.circle` function is used;
- `...`: further graphical parameters used to plot the graphs. See package `igraph` for more details.

**Details**

For a given value of the tuning parameter, specified by the argument `rhoid`, `gplot.fglasso` shows the graphs estimated at the time points `tp[1]` and `tp[2]`. By convention, the graph associated to the sub matrix \( K_{tp[1],tp[2]} \) is represented by a directed graph where a directed edge is drawn by an arrow from a vertex in the first graph pointing forwards a vertex in the second graph.
Value

gplot.fglasso returns a list with components:

- `graph.tp1`: an object with class `igraph` representing the undirected graph estimated at the time point `tp[1]`;
- `graph.tp2`: an object with class `igraph` representing the undirected graph estimated at the time point `tp[2]`;
- `graph.net`: an object with class `igraph` representing the directed graph associated to the submatrix $K_{tp[1],tp[2]}$;
- `layout`: the matrix used to specify the placement of the vertices.

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See Also

`fglasso` function.

Examples

```r
n <- 50
tp <- 3
p <- 3
X <- matrix(rnorm(n * p * tp), N, tp * p)
S <- crossprod(X) / N
model <- list(lag0 = c(s = "c", n = "ut"), lag1 = c(s = "t", n = "t"))
out.fglasso <- fglasso(S = S, model = model, tp = tp, p = p)
gplot(out.fglasso, rhoid = 50, sub.tp1 = "First graph", 
      sub.tp2 = "Second graph")
```

Description

`gplot.sglasso` shows the sequence of graphs estimated by `sglasso`.

Usage

```r
## S3 method for class 'sglasso'
gplot(object, rhoid, layout = layout.circle, ...)
```
**Arguments**

- **object**: fitted sglasso object;
- **rhoid**: vector of integers used to specify the $\rho$-values used to fit the sglasso model. By default `gplot.sglasso` shows the sequence of graphs estimated by `sglasso`. Only topologically different graphs are plotted;
- **layout**: a function or a matrix used to specify the layout of the graphs that will be plotted. By default the `layout.circle` function is used;
- **...** further graphical parameters used to plot the graphs. See package `igraph` for more details.

**Details**

gplot.sglasso shows the sequence of topologically different graphs estimated by `sglasso`. To specify the layout of the graphs, the user can use any layout function available in the R package `igraph`. The user can also specify the placement of the vertices by a matrix with two columns and the same number of rows as the number of vertices.

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**See Also**

- `sglasso` function.

**Examples**

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X)/N
mask <= outer(1:p, 1:p, function(i,j) 0.5-abs(i-j))
out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
gplot(out.sglasso_path)
gplot(out.sglasso_path, rhoid = 1:5)
```

**Description**

Function `Kh` computes the sequence of sparse structured precision matrices estimated by `sglasso` function.
Usage

\texttt{kh(object, rho)}

Arguments

- \texttt{object}: fitted \texttt{sglasso} object;
- \texttt{rho}: a subset of the values of the tuning parameter used in \texttt{sglasso} to compute the solution path. By default, the entire sequence of estimated sparse structured precision matrices is returned.

Value

\texttt{kh} returns a named list containing the sequence of estimated sparse structured precision matrices.

Author(s)

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See Also

\texttt{sglasso} function.

Examples

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i, j) 0.5^abs(i - j))
out.sglasso_path <- sglasso(S, mask, nrho = 5, tol = 1.0e-13)
out.sglasso_path
kh(out.sglasso_path)
rho <- out.sglasso_path$rho[3]
out.sglasso_single <- sglasso(S, mask, nrho = 1, min_rho = rho,
                           tol = 1.0e-13, algorithm = "ccm")
kh(out.sglasso_single)
```

---

\textbf{klcv} \hspace{1cm} \textit{Cross-Validated Kullback-Leibler Divergence}

Description

Model selection criterion based on the leave-one-out cross-validated Kullback-Leibler divergence.

Usage

\texttt{klcv(object, X, scale = 1)}
**Arguments**

- `object`: fitted `sglasso`/`fglasso` object;
- `X`: the matrix used to compute the empirical variance/covariance matrix. Its dimension is \( N \times p \), where \( p \) is the number of random variables and \( N \) is the sample size;
- `scale`: scalar value used to scale the estimated degrees-of-freedom. See below for more details.

**Details**

`klcv` function implements the leave-one-out cross-validate Kullback-Leibler divergence criterion proposed in Vujacic et al. (2015). For \( l_1 \)-penalized Gaussian Graphical Models this measure of goodness-of-fit has the following form

\[
klcv(\rho) = -\frac{\ell(\hat{K}(\rho))}{N} + \frac{\text{scale}}{2N} gdf(\hat{K}(\rho)),
\]

where \( \hat{K}(\rho) \) is the glasso estimate of the concentration matrix, \( \ell(\hat{K}(\rho)) \) is the corresponding value of the log-likelihood function, `scale` is a scale factor for the complexity part, i.e. \( gdf(\hat{K}(\rho)) \), which is defined as

\[
gdf(\hat{K}(\rho)) = \frac{1}{N-1} \sum_{k=1}^{N} vec\{(\hat{K}(\rho)^{-1} - S_k) \circ 1_{\rho}\} vec[\hat{K}(\rho)\{(S - S_k) \circ 1_{\rho}\} \hat{K}(\rho)].
\]

In the previous expression \( S \) is the empirical variance/covariance matrix, \( S_k = X_k X_k' \), \( 1_{\rho} \) is a matrix with entries \( I(k_{ij}(\rho) \neq 0) \) and \( \circ \) is the Hadamard product operator.

**Value**

`klcv` returns an S3 object with calls `klcv`, i.e. a named list with the following components:

- `klcv`: the vector with the leave-one-out cross-validated Kullback-Leibler divergence;
- `rho`: the rho-values used to compute the leave-one-out cross-validated Kullback-Leibler divergence;
- `loglik`: a vector with the log-likelihood computed for the sequence of weighted \( l_1 \)-penalized \( RCON(V, E) \);
- `gdf`: a vector returning the generalized degrees-of-freedom;
- `scale`: the scale value used to define the leave-one-out cross-validated Kullback-Leibler divergence;
- `min.klcv`: minimum value of the leave-one-out cross-validated Kullback-Leibler divergence;
- `rho.opt`: the rho-value corresponding to minimum leave-one-out cross-validated Kullback-Leibler divergence;
- `rhoid`: the index of the rho-value identified by the leave-one-out cross-validated Kullback-Leibler divergence.
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References


See Also

*sglasso*, *loglik* functions and *plot.klcv* method.

Examples

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i, j) 0.5^abs(i-j))
out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
out.klcv <- klcv(out.sglasso_path, X)
out.klcv
```

---

**loglik**  
*Extract Log-Likelihood*

Description

This function extracts the log-likelihood for the sequence of weighted l1-penalized $R CON(V, E)$ models estimated by *sglasso* function.

Usage

```r
loglik(object, N = 2)
```

Arguments

- `object` a fitted *sglasso* object;
- `N` sample size. Default value is 2 to remove the constant term in the log-likelihood function. See below for more details.
**Details**

Denoted with $\psi = (\eta', \theta')'$ the parameter vector of the structured concentration matrix $K(\psi)$, the log-likelihood function of the RCON($V, E$) model is equal, up to a constant, to the following expression

$$\ell(\psi) = \frac{N}{2} \left[ \log \det K(\psi) - tr \{SK(\psi) \} \right],$$

where $S = N^{-1} \sum_{i=1}^{N} X_i X_i^T$, $N$ is the sample size and $X_i$ is the $i$th observed $p$-dimensional vector. Denoted with $\hat{\psi} = (\hat{\eta}', \hat{\theta}')'$ the sglasso estimates, straightforward algebra shows that

$$\ell(\hat{\psi}) = \frac{N}{2} \left[ \log \det K(\hat{\psi}) - p + \rho \sum_{m=1}^{s} w_m |\hat{\theta}_m| \right],$$

where $\rho$ is the tuning parameter and $w_m$ are the weights used to define the weighted $l_1$-norm.

**Value**

loglik returns a vector containing the log-likelihood computed for the sequence of weighted $l_1$-penalized RCON($V, E$).

**Author(s)**

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**See Also**

summary.sglasso method and sglasso function.

**Examples**

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i, j) 0.5*abs(i-j))
out.sglasso_path <- sglasso(S, mask, nrho = 5, tol = 1.0e-13)
out.sglasso_path
loglik(out.sglasso_path, N = N)
rho <- out.sglasso_path$rho[3]
out.sglasso_single <- sglasso(S, mask, nrho = 1, min_rho = rho,
   tol = 1.0e-13, algorithm = "cvm")
loglik(out.sglasso_single, N = N)
```
neisseria

Description
This data set contains the gene expression data from a high-resolution time-course experiment based on the sequenced Neisseria meningitidis serogroup strain B strain MC58. Specifically, the expression level of ten genes is measured at ten different time points. Each column is standardized to have zero mean and standard deviation equal to one.

Usage
data("neisseria")

plot.klcv

Plot Method for Leave-One-Out Cross-Validated Kullback-Leibler Divergence

Description
plot.klcv produces a plot to study the sequence of leave-one-out cross-validated Kullback-Leibler divergences computed by klcv.

Usage
## S3 method for class 'klcv'
plot(x, ...)

Arguments

\begin{itemize}
  \item \textbf{x} \hspace{1cm} fitted klcv object;
  \item \textbf{...} \hspace{1cm} other parameters to be passed through the plotting function.
\end{itemize}

Details
This method function produces a plot showing the sequence of leave-one-out cross-validated Kullback-Leibler as function of the tuning parameter \textit{rho}. The optimal value of the tuning parameter is identified by a vertical dashed line.

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See Also

klcv function.

Examples

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i, j) 0.5^abs(i-j))
out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
out.klcv <- klcv(out.sglasso_path, X)
plot(out.klcv)
```

plot.sglasso

Plot Method for the Weighted L1-Penalized RCON(V, E) Model

Description

plot.sglasso produces two plots to study the sequence of models estimates by sglasso or fglasso.

Usage

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

Arguments

- `x` fitted sglasso/fglasso object;
- `...` other parameters to be passed through the plotting function.

Details

This function produces two different plots. The first one shows the path of the estimated parameters as function of the tuning parameter \( \rho \). In the same way, the second plot shows the path of the weighted scores as function of \( \rho \).

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See Also

sglasso function and summary.sglasso method.
Examples

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X)/N
mask <- outer(1:p, 1:p, function(i, j) 0.5^abs(i-j))
mask[5,] <- mask[4,] <- mask[2,] <- NA
out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
plot(out.sglasso_path)
```

sglasso  

**Lasso Method for the RCON(V,E) Models**

**Description**

Fit the weighted l1-penalized RCON(V,E) models using a cyclic coordinate algorithm.

**Usage**

```r
sglasso(S, mask, w = NULL, flg = NULL, min_rho = 1.0e-02, nrho = 50,
nstep = 1.0e+05, algorithm = c("ccd","ccm"), truncate = 1e-05,
       tol = 1.0e-03)
```

**Arguments**

- `S`: the empirical variance/covariance matrix;
- `mask`: a symmetric matrix used to specify the equality constraints on the entries of the concentration matrix. See the example below for more details;
- `w`: a vector specifying the weights used to compute the weighted l1-norm of the parameters of the RCON(V,E) model;
- `flg`: a logical vector used to specify if a parameter is penalized, i.e., if `flg[i] = TRUE` then the i-th parameter is penalized, otherwise (`flg[i] = FALSE`) the maximum likelihood estimate is computed;
- `min_rho`: last value of the sequence of tuning parameters used to compute the sglasso solution path. If `nrho = 1`, then `min_rho` is the value used to compute the sglasso estimate. Default value is 1.0e-02;
- `nrho`: number of tuning parameters used to compute the sglasso solution path. Default is 50;
- `nstep`: nonnegative integer used to specify the maximum number of iterations of the two cyclic coordinate algorithms. Default is 1.0e+05;
- `algorithm`: character by means of to specify the algorithm used to fit the model, i.e., a cyclic coordinate descente (ccd) algorithm or a cyclic coordinate minimization (ccm) algorithm. Default is ccd;
- `truncate`: at convergence all estimates below this value will be set to zero. Default is 1e-05;
- `tol`: value used for convergence. Default value is 1.0e-05.
Details

The RCON(V, E) model (Hojsgaard et al., 2008) is a kind of restriction of the Gaussian Graphical Model defined using a coloured graph to specify a set of equality constraints on the entries of the concentration matrix. Roughly speaking, a coloured graph implies a partition of the vertex set into \( R \) disjoint subsets, called vertex colour classes, and a partition of the edge set into \( S \) disjoint subsets, called edge colour classes. At each vertex/edge colour class is associated a specific colour. If we denote by \( K = (k_{ij}) \) the concentration matrix, i.e. the inverse of the variance/covariance matrix \( \Sigma \), the coloured graph implies the following equality constraints:

1. \( k_{ii} = \eta_n \) for any index \( i \) belonging to the \( n \)th vertex colour class;
2. \( k_{ij} = \theta_m \) for any pair \((i, j)\) belonging to the \( m \)th edge colour class.

Denoted with \( \psi = (\eta', \theta')' \) the \((R+S)\)-dimensional parameter vector, the concentration matrix can be defined as

\[
K(\psi) = \sum_{n=1}^{R} \eta_n D_n + \sum_{m=1}^{S} \theta_m T_m,
\]

where \( D_n \) is a diagonal matrix with entries \( D^n_{ii} = 1 \) if the index \( i \) belongs to the \( n \)th vertex colour class and zero otherwise. In the same way, \( T_m \) is a symmetric matrix with entries \( T^m_{ij} = 1 \) if the pair \((i, j)\) belongs to the \( m \)th edge colour class. Using the previous specification of the concentration matrix, the structured graphical lasso (sglasso) estimator (Abbruzzo et al., 2014) is defined as

\[
\hat{\psi} = \arg \max_{\psi} \log \det K(\psi) - \text{tr}\{S k(\psi)\} - \rho \sum_{m=1}^{S} w_m |\theta_m|,
\]

where \( S \) is the empirical variance/covariance matrix, \( \rho \) is the tuning parameter used to control the amount of shrinkage and \( w_m \) are weights used to define the weighted \( \ell_1 \)-norm. By default, the sglasso function sets the weights equal to the cardinality of the edge colour classes.

Value

sglasso returns an object with S3 class "sglasso", i.e. a named list containing the following components:

call the call that produced this object;
	nv number of vertex colour classes;

tne number of edge colour classes;

theta the matrix of the sglasso estimates. The first \( nv \) rows correspond to the unpenalized parameters while the remaining rows correspond to the weighted \( \ell_1 \)-penalized parameters;

w the vector of weights used to define the weighted \( \ell_1 \)-norm;

df \( \nu \rho \)-dimensional vector of the number of estimated nonzero parameters;

rho \( \nu \rho \)-dimensional vector of the sequence of tuning parameters;

grd the matrix of the scores;

nstep nonnegative integer used to specify the maximum number of iterations of the algorithms;
number of tuning parameters used to compute the sglasso solution path;
the algorithm used to fit the model;
the value used to set to zero the estimated parameters;
a nonnegative value used to define the convergence of the algorithms;
the empirical variance/covariance matrix used to compute the sglasso solution path;
the mask used to define the equality constraints on the entries of the concentration matrix;
number of iterations of the algorithm;
an integer value used to encode the warnings related to the algorithms. If conv = 0 the convergence has been achieved otherwise the maximum number of iterations has been achieved.

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References


See Also
summary.sglasso, plot.sglasso gplot.sglasso and methods.
The function Kh extracts the estimated sparse structured concentration matrices.

Examples
########################################################################
# sglasso solution path
#
## structural zeros:
## there are two ways to specify structural zeros which are
## related to the kind of mask. If mask is a numeric matrix
## NA is used to identify the structural zero. If mask is a
## character matrix then the structural zeros are specified
## using NA or ".".
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i,j) 0.5*abs(i-j))
mask

out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
out.sglasso_path

rho <- out.sglasso_path$rho[20]
out.sglasso <- sglasso(S, mask, nrho = 1, min_rho = rho, tol = 1.0e-13, algorithm = "ccm")
out.sglasso

out.sglasso_path$theta[, 20]
out.sglasso$theta[, 1]

summary.sglasso

**Summarizing sglasso Fits**

**Description**

summary method for class "sglasso".

**Usage**

```r
## S3 method for class 'sglasso'
summary(object, N, k = c("bic", "aic"),
         digits = max(3, getOption("digits") - 3), ...)
```

**Arguments**

- `object` fitted sglasso object;
- `N` sample size;
- `k` character/numeric argument used to specify the 'weight' of the complexity part in the measure of goodness-of-fit used to select the best model (see below for more details). Default is `k = "bic"`;
- `digits` significant digits in printout;
- `...` additional print arguments.

**Details**

`summary.sglasso` gives us information about the sequence of models estimated by the sglasso estimator. To select the best model, summary method uses a measure of Goodness-of-Fit (GoF) defined as follows:

\[-2\ell(\hat{\psi}) + k \times df,\]

where \( \ell(\hat{\psi}) \) is the log-likelihood of the estimated weighted l1-penalized \( RCON(V, E) \) model, \( df \) is the number of nonzero estimated parameters and \( k \) is a non-negative value used to weight the complexity part in the measure of goodness-of-fit. By default the summary method computes the BIC criterion to select the best model (\( k = "bic" \)). The AIC criterion can be easily computed.
The summary method is divided in two sections. First section shows the call producing the argument object followed by a data.frame. The column named \( \rho \) shows the sequence of the \( \rho \) values used to compute the solution curve, while the column log-lik shows the corresponding values of the log-likelihood function. The remaining columns show the number of estimated non-zero parameters, the values of the GoF and the associated ranking of the estimated models. Finally, the second section shows the estimated parameters of the best model identified by the used GoF criterion. Informations about the algorithm and the corresponding convergence are also provided.

Value

A list with components table and theta_gof is silently returned. The table component is the data.frame previously described while the component theta_gof is the vector of the estimated parameters corresponding to the best models identified by the GoF criterion.

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See Also

sglasso and loglik functions.

Examples

```r
N <- 100
p <- 5
X <- matrix(rnorm(N * p), N, p)
S <- crossprod(X) / N
mask <- outer(1:p, 1:p, function(i, j) 0.5^abs(i-j))
out.sglasso_path <- sglasso(S, mask, tol = 1.0e-13)
summary(out.sglasso_path, N)
rho <- out.sglasso_path$rho[20]
out.sglasso <- sglasso(S, mask, nrho = 1, min_rho = rho, tol = 1.0e-13)
summary(out.sglasso, N)
```
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