# Package 'GGMridge'

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Type Package
<b>Title</b> Gaussian Graphical Models Using Ridge Penalty Followed by Thresholding and Reestimation
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Depends mytnorm, MASS, stats, graphics
<b>Description</b> Estimation of partial correlation matrix using ridge penalty followed by thresholding and reestimation. Under multivariate Gaussian assumption, the matrix constitutes an Gaussian graphical model (GGM).
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R topics documented:
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## Description

Estimation of the parameters, null proportion, and degrees of freedom of the exact null density in the mixture distribution.

#### Usage

```
EM.mixture(p, eta0, df, tol)
```

## **Arguments**

p	A numeric vector representing partial correlation coefficients.
eta0	An initial value for the null proportion; 1-eta0 is the non-null proportion.
df	An initial value for the degrees of freedom of the exact null density.
tol	The tolerance level for convergence.

#### Value

## A list object containing

df Estimated degrees of freedom of the null density.

eta0 Estimated null proportion.

iter The number of iterations required to reach convergence.

#### Author(s)

Min Jin Ha

### References

Schafer, J. and Strimmer, K. (2005). An empirical Bayes approach to inferring large-scale gene association networks. Bioinformatics, 21, 754–764.

getEfronp 3

getEfronp	Estimation of empirical null distribution.	
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## Description

Estimation of empirical null distribution using Efron's central matching.

## Usage

```
getEfronp(
  z,
  bins = 120L,
  maxQ = 9,
  pct = 0,
  pct0 = 0.25,
  cc = 1.2,
  plotIt = FALSE
)
```

## Arguments

Z	A numeric vector of z values following the theoretical normal null distribution.
bins	The number of intervals for density estimation of the marginal density of z.
maxQ	The maximum degree of the polynomial to be considered for density estimation of the marginal density of z.
pct	Low and top ( $pct*100$ ) $f(z)$ .
pct0	Low and top (pct $0*100$ ) estimate f0(z).
СС	The central parts
	$(\mu - \sigma cc, \mu + \sigma cc)$
	of the empirical distribution $z$ are used for an estimate of the null proportion (eta).
plotIt	TRUE if density plot is to be produced.

## Value

## A list containing

correctz	The corrected z values to follow empirically standard normal distribution.
correctp	The corrected p values using the correct z values.
q	The chosen degree of polynomial for the estimated marginal density.
mu0hat	The location parameter for the normal null distribution.
sigma0hat	The scale parameter for the normal null distribution.
eta	The estimated null proportion.

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#### Author(s)

Min Jin Ha

#### References

Efron, B. (2004). Large-scale simultaneous hypothesis testing. Journal of the American Statistical Association, 99, 96–104.

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

#### **Examples**

```
p <- 100 # number of variables
n <- 50 # sample size
#####################################
# Simulate data
simulation \leftarrow simulateData(G = p, etaA = 0.02, n = n, r = 1)
data <- simulation$data[[1]]</pre>
stddata <- scale(x = data, center = TRUE, scale = TRUE)</pre>
#####################################
# estimate ridge parameter
lambda.array <- seq(from = 0.1, to = 20, by = 0.1) * (n - 1.0)
fit <- lambda.cv(x = stddata, lambda = lambda.array, fold = 10L)</pre>
lambda <- fit$lambda[which.min(fit$spe)] / (n - 1.0)</pre>
###################################
# calculate partial correlation
# using ridge inverse
w.upper <- which(upper.tri(diag(p)))</pre>
partial <- solve(lambda * diag(p) + cor(data))</pre>
partial <- (-scaledMat(x = partial))[w.upper]</pre>
# get p-values from empirical
# null distribution
efron.fit <- getEfronp(z = transFisher(x = partial))</pre>
```

ksStat

The Kolmogorov-Smirnov Statistic for p-Values

#### **Description**

Calculates the Kolmogorov-Smirnov statistic for p-values

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## Usage

```
ksStat(p)
```

#### **Arguments**

p A numeric vector with p-values.

#### Value

Kolmogorov-Smirnov statistic

#### Author(s)

Min Jin Ha

## **Examples**

```
p <- stats::runif(100)
ksStat(p = p)
ks.test(p, y = "punif") # compare with ks.test</pre>
```

lambda.cv

Choose the Tuning Parameter of the Ridge Inverse

#### **Description**

Choose the tuning parameter of the ridge inverse by minimizing cross validation estimates of the total prediction errors of the p separate ridge regressions.

## Usage

```
lambda.cv(x, lambda, fold)
```

#### **Arguments**

x An n by p data matrix.

lambda A numeric vector of candidate tuning parameters.

fold fold-cross validation is performed.

## Value

A list containing

lambda The selected tuning parameter, which minimizes the total prediction errors.

spe The total prediction error for all the candidate lambda values.

6 lambda.pcut.cv

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

#### **Examples**

```
p <- 100 # number of variables
n <- 50 # sample size
#####################################
# Simulate data
#####################################
simulation \leftarrow simulateData(G = p, etaA = 0.02, n = n, r = 1)
data <- simulation$data[[1L]]</pre>
stddata <- scale(x = data, center = TRUE, scale = TRUE)</pre>
###################################
# estimate ridge parameter
lambda.array <- seq(from = 0.1, to = 20, by = 0.1) * (n - 1.0)
fit <- lambda.cv(x = stddata, lambda = lambda.array, fold = 10L)</pre>
lambda <- fit$lambda[which.min(fit$spe)] / (n - 1.0)</pre>
# calculate partial correlation
# using ridge inverse
partial <- solve(lambda*diag(p) + cor(data))</pre>
partial <- -scaledMat(x = partial)</pre>
```

lambda.pcut.cv

Choose the Tuning Parameter of the Ridge Inverse and Thresholding Level of the Empirical p-Values

#### **Description**

Choose the tuning parameter of the ridge inverse and p-value cutoff by minimizing cross validation estimates of the total prediction errors of the p separate ridge regressions.

### Usage

```
lambda.pcut.cv(x, lambda, pcut, fold = 10L)
```

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## Arguments

x n by p data matrix.

lambda A vector of candidate tuning parameters.pcut A vector of candidate cutoffs of pvalues.

fold fold-cross validation is performed.

#### Value

The total prediction errors for all lambda (row-wise) and pcut (column-wise)

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

```
p <- 100 # number of variables
n <- 50 # sample size
# Simulate data
simulation \leftarrow simulateData(G = p, etaA = 0.02, n = n, r = 1)
data <- simulation$data[[1L]]</pre>
stddata <- scale(x = data, center = TRUE, scale = TRUE)</pre>
# Selection of a lambda and a
# p-value cutoff
#####################################
lambda.array \leftarrow seq(from = 0.1, to = 5, length = 10) * (n-1.0)
pcut.array <- seq(from = 0.01, to = 0.05, by = 0.01)
tpe <- lambda.pcut.cv(x = stddata,
                     lambda = lambda.array,
                     pcut = pcut.array,
                     fold = 3L)
w.mintpe <- which(tpe == min(tpe), arr.ind = TRUE)</pre>
lambda <- lambda.array[w.mintpe[1L]]</pre>
alpha <- pcut.array[w.mintpe[2L]]</pre>
```

8 lambda.pcut.cv1

lambda.pcut.cv1	Choose the Tuning Parameter of the Ridge Inverse and Thresholding Level of the Empirical p-Values. Calculate total prediction error for test data after fitting partial correlations from train data for all values of lambda and pcut.
	of lambda and pcut.

#### **Description**

Choose the Tuning Parameter of the Ridge Inverse and Thresholding Level of the Empirical p-Values.

Calculate total prediction error for test data after fitting partial correlations from train data for all values of lambda and pcut.

#### Usage

```
lambda.pcut.cv1(train, test, lambda, pcut)
```

#### **Arguments**

train	An n x p data matrix from which the model is fitted.
test	An m x p data matrix from which the model is evaluated.
lambda	A vector of candidate tuning parameters.
pcut	A vector of candidate cutoffs of pyalues.

#### Value

Total prediction error for all the candidate lambda and pvalue cutoff values.

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

lambda.TargetD

```
####################################
# Split into train/test sets
####################################
testindex <- sample(1L:n, 10L)</pre>
train <- data[-testindex,,drop = FALSE]</pre>
stdTrain <- scale(x = train, center = TRUE, scale = TRUE)</pre>
test <- data[testindex,,drop = FALSE]</pre>
stdTest <- scale(x = test, center = TRUE, scale = TRUE)</pre>
####################################
# Calculate total prediction
# errors for all candidate
# lambda and p-value cutoffs
lambda.array <- seq(from = 0.1, to = 5, length = 10) * (n - 1.0)
pcut.array <- seq(from = 0.01, to = 0.05, by = 0.01)
tpe <- lambda.pcut.cv1(train = stdTrain,</pre>
                        test = stdTest,
                        lambda = lambda.array,
                        pcut = pcut.array)
```

lambda.TargetD

Shrinkage Estimation of a Covariance Matrix Toward an Identity Matrix

## **Description**

Estimation of a weighted average of a sample covariance (correlation) matrix and an identity matrix.

#### Usage

```
lambda.TargetD(x)
```

#### **Arguments**

x Centered data for covariance shrinkage and standardized data for correlation shrinkage.

### **Details**

An analytical approach to the estimate ridge parameter.

#### Value

The estimates of shrinkage intensity.

10 ne.lambda.cv

#### Author(s)

Min Jin Ha

#### References

Schafer, J. and Strimmer, K. (2005). A shrinkage approach to large-scale covariance matrix estimation and implications for functional genomics. Statistical Applications in Genetics and Molecular Biology, 4, 32.

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

#### **Examples**

```
##################################
# Simulate data
simulation <- simulateData(G = 100, etaA = 0.02, n = 50, r = 10)
dat <- simulation$data[[1L]]</pre>
stddat <- scale(x = dat, center = TRUE, scale = TRUE)</pre>
shrinkage.lambda <- lambda.TargetD(x = stddat)</pre>
# the ridge parameter
####################################
ridge.lambda <- shrinkage.lambda / (1.0 - shrinkage.lambda)</pre>
# partial correlation matrix
####################################
partial <- solve(cor(dat) + ridge.lambda * diag(ncol(dat)))</pre>
partial <- -scaledMat(x = partial)</pre>
```

ne.lambda.cv

Choose the Tuning Parameter of a Ridge Regression Using Cross-Validation

#### **Description**

Choose the tuning parameter of a ridge regression using cross-validation.

## Usage

```
ne.lambda.cv(y, x, lambda, fold)
```

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## Arguments

y Length n response vector.

x n x p matrix for covariates with p variables and n sample size.

lambda A numeric vector for candidate tuning parameters for a ridge regression.

fold fold-cross validation used to choose the tuning parameter.

#### Value

A list containing

1 The selected tuning parameter, which minimizes the prediction error.

spe The prediction error for all of the candidate lambda values.

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

12 R.separate.ridge

R.separate.ridge	Estimation of Partial Correlation Matrix Using p Separate Ridge Regressions.
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#### **Description**

The partial correlation matrix is estimated by p separate ridge regressions with the parameters selected by cross validation.

#### Usage

```
R.separate.ridge(x, fold, lambda, verbose = FALSE)
```

#### **Arguments**

x n x p data matrix; n is the # of samples and p is the # of variables.

fold Ridge parameters are selected by fold-cross validations separately for each re-

gression.

lambda The candidate ridge parameters for all p ridge regressions.

verbose TRUE/FALSE; if TRUE, print the procedure.

#### Value

A list containing

R The partial correlation matrix.

lambda.sel The selected tuning parameters for p ridge regressions.

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

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scaledMat

Scale a square matrix

#### **Description**

Scale a square matrix to have unit diagonal elements.

#### Usage

```
scaledMat(x)
```

#### **Arguments**

х

A square matrix with positive diagonal elements

#### Value

Scaled matrix of x.

## Author(s)

Min Jin Ha

14 simulateData

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Generate Simulation Data from a Random Network.

#### **Description**

Generate a random network where both the network structure and the partial correlation coefficients are random. The data matrices are generated from multivariate normal distribution with the covariance matrix corresponding to the network.

#### Usage

```
simulateData(G, etaA, n, r, dist = "mvnorm")
```

#### **Arguments**

G	The number of	variables (	(vertices).

etaA The proportion of non-null edges among all the G(G-1)/2 edges.

n The sample size.

r The number of replicated G by N data matrices.

dist A function which indicates the distribution of sample. "mvnorm" is multivari-

ate normal distribution and "mvt" is multivariate t distribution with df=2. The

default is set by "mvnorm".

#### Value

A list containing

data a list, each element containing an n X G matrix of simulated data.

true.partialcor

The partial correlation matrix which the datasets are generated from.

truecor.scaled

The covariance matrix calculted from the partial correlation matrix.

sig. node The indices of nonzero upper triangle elements of partial correlation matrix.

## Author(s)

Min Jin Ha

#### References

Schafer, J. and Strimmer, K. (2005). An empirical Bayes approach to inferring large-scale gene association networks. Bioinformatics, 21, 754–764.

```
simulation <- simulateData(G = 100, etaA = 0.02, n = 50, r = 10)
```

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structuredEstimate

Estimation of Partial Correlation Matrix Given Zero Structure.

#### **Description**

Estimation of nonzero entries of the partial correlation matrix given zero structure.

#### Usage

```
structuredEstimate(x, E)
```

#### **Arguments**

x n by p data matrix with the number of variables p and sample size n.

E The row and column indices of zero entries of the partial correlation matrix.

#### Value

A list containing

R The partial correlation matrix.
K The inverse covariance matrix.
RSS The residual sum of squares.

#### Author(s)

Min Jin Ha

#### References

Ha, M. J. and Sun, W. (2014). Partial correlation matrix estimation using ridge penalty followed by thresholding and re-estimation. Biometrics, 70, 762–770.

16 transFisher

```
lambda.array <- seq(from = 0.1, to = 20, by = 0.1) * (n-1.0)
fit <- lambda.cv(x = stddata, lambda = lambda.array, fold = 10L)</pre>
lambda <- fit$lambda[which.min(fit$spe)]/(n-1)</pre>
# calculate partial correlation
# using ridge inverse
####################################
w.upper <- which(upper.tri(diag(p)))</pre>
partial <- solve(lambda * diag(p) + cor(data))</pre>
partial <- (-scaledMat(x = partial))[w.upper]</pre>
####################################
# get p-values from empirical
# null distribution
efron.fit <- getEfronp(z = transFisher(x = partial),</pre>
                     bins = 50L,
                     maxQ = 13)
# estimate the edge set of
# partial correlation graph with
# FDR control at level 0.01
####################################
w.array <- which(upper.tri(diag(p)),arr.ind=TRUE)</pre>
th <- 0.01
wsig <- which(p.adjust(efron.fit$correctp, method="BH") < th )</pre>
E <- w.array[wsig,]</pre>
dim(E)
# structured estimation
fit <- structuredEstimate(x = stddata, E = E)</pre>
th.partial <- fit$R</pre>
```

transFisher

Fisher's Z-Transformation

#### Description

Fisher's Z-transformation of (partial) correlation.

#### Usage

```
transFisher(x)
```

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#### **Arguments**

A vector having entries between -1 and 1.

#### Value

Fisher's Z-transformed values.

#### Author(s)

Min Jin Ha

```
# Simulate data
simulation <- simulateData(G = 100, etaA = 0.02, n = 50, r = 1)
dat <- simulation$data[[1L]]</pre>
stddat <- scale(x = dat, center = TRUE, scale = TRUE)</pre>
shrinkage.lambda <- lambda.TargetD(x = stddat)</pre>
# the ridge parameter
ridge.lambda <- shrinkage.lambda / (1.0 - shrinkage.lambda)</pre>
# partial correlation matrix
partial <- solve(cor(dat) + ridge.lambda * diag(ncol(dat)))</pre>
partial <- -scaledMat(x = partial)</pre>
# Fisher's Z transformation of
# upper diagonal of the partial
# correlation matrix
####################################
w.upper <- which(upper.tri(diag(nrow(dat))))</pre>
psi <- transFisher(x = partial[w.upper])</pre>
```

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