

Package ‘heterocop’

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Type Package

Title Semi-Parametric Estimation with Gaussian Copula

Version 0.1.0.0

Description A method for generating random vectors which are linked by a Gaussian copula. It also enables to estimate the correlation matrix of the Gaussian copula in order to identify independencies within the data.

Imports mvtnorm, stats, igraph, matrixcalc, graphics, foreach,
stringr, doSNOW, utils

License GPL (>= 3)

Encoding UTF-8

LazyData true

RoxygenNote 7.3.2

Depends R (>= 2.10)

Suggests knitr, rmarkdown, kableExtra, dplyr

VignetteBuilder knitr

NeedsCompilation no

Author Julie Cartier [aut],
Florence Jaffrezic [aut],
Gildas Mazo [aut],
Ekaterina Tomilina [aut, cre]

Maintainer Ekaterina Tomilina <ekaterina.tomilina@inrae.fr>

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CopulaSim

*CopulaSim***Description**

This function enables the user to simulate data from a Gaussian copula and arbitrary marginal quantile functions

Usage

```
CopulaSim(n, R, qdist, random = FALSE)
```

Arguments

- n the number of observations
- R a correlation matrix of size dxd
- qdist a vector containing the names of the marginal quantile functions as well as the number of times they are present in the dataset
- random a boolean defining whether the order of the correlation coefficients should be randomized

Value

a list containing an nxn data frame, the shuffled correlation matrix R, and the permutation leading to the new correlation matrix

Examples

```
M <- diag_block_matrix(c(3,4,5),c(0.7,0.8,0.2))
CopulaSim(20,M,c(rep("qnorm(0,1)",6),rep("qexp(0.5)",4),rep("qbinom(4,0.8)",2)),random=TRUE)
```

cor_network_graph

cor_network_graph

Description

This function enables the user to plot the graph corresponding to the correlations of the Gaussian copula

Usage

```
cor_network_graph(R, TS, binary = TRUE, legend)
```

Arguments

R	a correlation matrix of size dxd (d is the number of variables)
TS	a threshold for the absolute values of the correlation matrix coefficients
binary	a boolean specifying whether the coefficients should be binarized, TRUE by default (zero if the coefficient is less than the threshold in absolute value, 1 otherwise). If FALSE, the edge width is proportional to the coefficient value.
legend	a vector containing the type of each variable used to color the vertices

Value

a graph representing the correlations between the latent Gaussian variables

Examples

```
R <- diag_block_matrix(c(3,4,5),c(0.7,0.8,0.2))
data <- CopulaSim(20,R,c(rep("qnorm(0,1)",6),rep("qexp(0.5)",4),
rep("qbinom(4,0.8)",2)),random=FALSE)[[1]]
cor_network_graph(R,TS=0.3,binary=TRUE,legend=c(rep("Normal",6),
rep("Exponential",4),rep("Binomial",2)))
```

diag_block_matrix

diag_block_matrix

Description

This function enables the user to generate a diagonal block-matrix with homogeneous blocks

Usage

```
diag_block_matrix(blocks, coeff)
```

Arguments

- blocks** a vector containing the sizes of the blocks
coeff a vector containing the coefficient corresponding to each block, the coefficients must be between 0 and 1

Value

a diagonal block-matrix containing the specified coefficients

Examples

```
diag_block_matrix(c(3,4,5),c(0.3,0.4,0.8))
```

gauss_gen

gauss_gen

Description

This function enables the user to generate gaussian vectors with correlation matrix R

Usage

```
gauss_gen(R, n)
```

Arguments

- R** a correlation matrix of size dxd
n the number of observations

Value

a nxn data frame containing n observations of the d variables

Examples

```
M <- diag_block_matrix(c(3,4,5),c(0.7,0.8,0.2))
gauss_gen(M,20)
```

icgc_data*ICGC dataset*

Description

Dataset containing RNA counts, protein expression and mutations measured on breast cancer tumors.

Usage

```
icgc_data
```

Format

A dataframe of 15 variables and 250 observations containing the following:

ACACA, AKT1S1, ANLN,ANXA1,AR RNA counts (discrete)

ACACA_P, AKT1S1_P, ANLN_P,ANXA_P,AR_P protein expression measurements (discrete)

MU5219,MU4468,MU7870,MU4842,MU6962 5 mutations (binary)

matrix_cor_ts*matrix_cor_ts*

Description

This function enables the user to threshold matrix coefficients

Usage

```
matrix_cor_ts(R, TS, binary = TRUE)
```

Arguments

R	a correlation matrix
TS	a threshold
binary	a boolean specifying whether the coefficients should be binarized, TRUE by default (zero if the coefficient is less than the threshold in absolute value, 1 otherwise)

Value

the thresholded input matrix

Examples

```
M <- diag_block_matrix(c(3,4,5),c(0.7,0.8,0.2))
matrix_cor_ts(M,0.5)
```

matrix_gen*matrix_gen***Description**

This function enables the user to generate a sparse, nonnegative definite correlation matrix via the Cholesky decomposition

Usage

```
matrix_gen(d, gamma)
```

Arguments

- | | |
|--------------|--|
| d | the number of variables |
| gamma | an initial sparsity parameter for the lower triangular matrices in the Cholesky decomposition, must be between 0 and 1 |

Value

a list containing the generated correlation matrix and its final sparsity parameter (ie the proportion of zeros)

Examples

```
matrix_gen(15, 0.81)
```

rho_estim*rho_estim***Description**

This function enables the user to estimate the correlation matrix of the Gaussian copula for a given dataset

Usage

```
rho_estim(data, Type, parallel = FALSE)
```

Arguments

- | | |
|-----------------|--|
| data | an nx d data frame containing n observations of d variables |
| Type | a vector containing the type of the variables, "C" for continuous and "D" for discrete |
| parallel | a boolean encoding whether the computations should be parallelized |

rho_estim

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Value

the dxd estimated correlation matrix of the Gaussian copula

Examples

```
M <- diag_block_matrix(c(3,4,5),c(0.7,0.8,0.2))
data <- CopulaSim(20,M,c(rep("qnorm(0,1)",6),rep("qexp(0.5)",4),
rep("qbinom(4,0.8)",2)),random=FALSE)[[1]]
rho_estim(data,c(rep("C",10),rep("D",2)))
```

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