Package 'singR'

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```
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Description Implementation of SING algorithm to extract joint and individual non-Gaussian compo-
      nents from two datasets. SING uses an objective function that maximizes the skewness and kur-
      tosis of latent components with a penalty to enhance the similarity between subject scores. Un-
      like other existing methods, SING does not use PCA for dimension reduc-
      tion, but rather uses non-Gaussianity, which can improve feature extraction. Ben-
      jamin B.Risk, Irina Gaynanova (2021) <doi:10.1214/21-AOAS1466>.
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```

Title Simultaneous Non-Gaussian Component Analysis

Type Package

2 angleMatchICA

R topics documented:

angleMatchICA Match the columns of Mx and My		Match the colums of Mx and My	
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Description

 $\label{eq:match_condition} \mbox{angleMatchICA match the colums of } Mx \mbox{ and } My, \mbox{ using the } n \mbox{ } x \mbox{ } p \mbox{ parameterization of the } JIN \mbox{ decomposition assumes}$

Usage

```
angleMatchICA(Mx, My, Sx = NULL, Sy = NULL)
```

Arguments

Mx	Subject score for X matrix of n x n.comp
My	Subject score for Y matrix of n x n.comp
Sx	Variable loadings for X matrix of n.comp x px
Sy	Variable loadings for Y matrix of n.comp x py

aveM 3

Value

a list of matrixes: ## Mx: ## My: ## matchedangles: ## allangles: ## perm: ## omangles:

aveM

Average Mj for Mx and My Here subjects are by rows, columns correspond to components

Description

Average Mj for Mx and My Here subjects are by rows, columns correspond to components

Usage

```
aveM(mjX, mjY)
```

Arguments

```
mjX n x rj mjY n x rj
```

Value

a new Mj

```
#get simulation data
data(exampledata)
data=exampledata

# To get n.comp value, we can use NG_number function.

# use JB statistic as the measure of nongaussianity to run lngca with df=0
output_JB=singR(dX=exampledata$dX,dY=exampledata$dY,
df=0,rho_extent="small",distribution="JB",individual=TRUE)

est.Mj = aveM(output_JB$est.Mjx,output_JB$est.Mjy)
```

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calculateJB Calculate determin	es the sum of the JB scores across all components, useful for ing rho.
--------------------------------	--

Description

We measure non-Gaussianity using Jarque-Bera (JB) statistic, which is a weighted combination of squared skewness and kurtosis, JB paper. The data has to be standardized and mean 0 and sd to 1.

Usage

```
calculateJB(S = NULL, U = NULL, X = NULL, alpha = 0.8)
```

Arguments

S the variable loadings r x px.

U matrix for matched columns rj x n

X whitened data matrix n x px, data = whitener XA %*% dXcentered

alpha JB weighting of skewness and kurtosis. default = 0.8

Value

the sum of JB score across all components.

covwhitener Returns square root of the precision matrix for whitening

Description

Returns square root of the precision matrix for whitening

Usage

```
covwhitener(X, n.comp = ncol(X), center.row = FALSE)
```

Arguments

X Matrix

n.comp the number of components

center.row whether to center

Value

square root of the precision matrix for whitening

create.graph.long 5

create.graph.long

create graph dataset with netmat and mmp_order a data.frame called with vectorization of reordered netmat by mmp_order.

Description

create graph dataset with netmat and mmp_order a data.frame called with vectorization of reordered netmat by mmp_order.

Usage

```
create.graph.long(gmatrix, sort_indices = NULL)
```

Arguments

```
gmatrix netmat
sort_indices mmp_order
```

Value

a data.frame with vectors: ## X1: vector of numerics. ## X2: vector of numerics. ## value: vectorization of reordered netmat by mmp_order.

curvilinear

Curvilinear algorithm with r0 joint components

Description

The curvilinear algorithm is modified from Wen and Yin paper.

Usage

```
curvilinear(
   Ux,
   Uy,
   xData,
   yData,
   invLx,
   invLy,
   rho,
   tau = 0.01,
   alpha = 0.8,
   maxiter = 1000,
   tol = 1e-06,
   rj
)
```

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Arguments

Ux Matrix with n.comp x n, initial value of Ux, comes from greedyMatch.

Uy Matrix with n.comp x n, initial value of Uy, comes from greedyMatch.

xData matrix with n x px, Xw = Lx %*% Xc. yData matrix with n x py, Yw = Ly %*% Yc. invLx Inverse matrix of Lx, matrix n x n. invLy Inverse matrix of Ly, matrix n x n.

rho the weight parameter of matching relative to non-gaussianity.

tau initial step size, default value is 0.01

alpha controls weighting of skewness and kurtosis. Default value is 0.8, which corre-

sponds to the Jarque-Bera test statistic with 0.8 weighting on squared skewness

and 0.2 on squared kurtosis.

maxiter default value is 1000

tol the threshold of change in Ux and Uy to stop the curvlinear function

rj the joint rank, comes from greedyMatch.

Value

a list of matrices:

Ux Optimized Ux with matrix n.comp x n.

Uy Optimized Uy with matrix $n.comp \ x \ n.$

tau step size

iter number of iterations.

error PMSE(Ux,Uxnew)+PMSE(Uy,Uynew)

obj Objective Function value

curvilinear_c

Curvilinear algorithm based on C code with r0 joint components

Description

#' The curvilinear algorithm is modified from Wen and Yin paper.

curvilinear_c 7

Usage

```
curvilinear_c(
   Ux,
   Uy,
   xData,
   yData,
   invLx,
   invLy,
   rho,
   tau = 0.01,
   alpha = 0.8,
   maxiter = 1000,
   tol = 1e-06,
   rj
)
```

Arguments

Ux	Matrix with n.comp x n, initial value of Ux, comes from greedyMatch.
Uy	Matrix with n.comp x n, initial value of Uy, comes from greedyMatch.
xData	matrix with n x px, $Xw = Lx \% *\% Xc$.
yData	matrix with n x py, $Yw = Ly \%*\% Yc$.
invLx	Inverse matrix of Lx, matrix n x n.
invLy	Inverse matrix of Ly, matrix n x n.
rho	the weight parameter of matching relative to non-gaussianity.
tau	initial step size, default value is 0.01
alpha	controls weighting of skewness and kurtosis. Default value is 0.8, which corresponds to the Jarque-Bera test statistic with 0.8 weighting on squared skewness and 0.2 on squared kurtosis.
maxiter	default value is 1000
tol	the threshold of change in Ux and Uy to stop the curvilinear function
rj	the joint rank, comes from greedyMatch.

Value

```
a list of matrices:

Ux Optimized Ux with matrix n.comp x n.

Uy Optimized Uy with matrix n.comp x n.

tau step size

iter number of iterations.

error PMSE(Ux,Uxnew)+PMSE(Uy,Uynew)

obj Objective Function value
```

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est.M.ols

Estimate mixing matrix from estimates of components

Description

Estimate mixing matrix from estimates of components

Usage

```
est.M.ols(sData, xData, intercept = TRUE)
```

Arguments

sData S rx x px xData dX n x px intercept default = TRUE

Value

a matrix Mx, dimension n x rx.

exampledata

Data for simulation example 1

Description

Data for simulation example 1

Usage

exampledata

Format

A data list with 10 subsets:

dX original data matrix for X with n x px, 48x3602

dY original data matrix for Y with n x py, 48x4950

mj true mj matrix, n x rj, 48x2

sIx true S matrix of independent non-Gaussian components in X, ri_x x px, 2x3602

sIy true S matrix of independent non-Gaussian components in Y, ri_y x py, 2x4950

sjx true S matrix of joint non-Gaussian components in X, rj x px, 2x3602

sjy true S matrix of joint non-Gaussian components in Y, rj x py, 2x4950

snr signal to noise ratio

R2x R2 for x data

R2y R2 for y data

gen.inits 9

gen.inits

Generate initialization from specific space

Description

Generate initialization from specific space

Usage

```
gen.inits(p, d, runs, orth.method = c("svd", "givens"))
```

Arguments

p p*p orthodox matrix
d p*d orthodox matrix

runs the number of orthodox matrix

orth.method orthodox method

Value

a list of initialization of mixing matrices.

Examples

```
gen.inits(2,3,3,'svd')
```

greedymatch

Greedy Match

Description

Greedy Match matches a column of Mx and My by minimizing chordal distance between vectors, removes the matched columns and then finds the next pair. This equivalent to maximizing absolute correlation for data in which each column has mean equal to zero. Returns permuted columns of Mx and My. This function does not do any scaling or sign flipping. For this matching to coincide with angle matching, the columns must have zero mean.

Usage

```
greedymatch(Mx, My, Ux, Uy)
```

10 lngca

Arguments

Mx	Subject Score for X with n x n.comp.X matrix
Му	Subject Score for Y with n x n.comp.Y matrix
Ux	Matrix with n.comp x n, $Mx = Lx^{-1} \% \% t Ux$, Lx is the whitener matrix of dX.
Uy	Matrix with n.comp x n, My = Ly $^-1$ %*% t Uy, Ly is the whitener matrix of dY.

Value

a list of matrices:

Mx Columns of original Mx reordered from highest to lowest correlation with matched component in My

My Columns of original My reordered from highest to lowest correlation with matched component in Mx

Ux Permuted rows of original Ux corresponds to MapX

Uy Permuted rows of original Uy corresponds to MapY

correlations a vector of correlations for each pair of columns in permuted Mx and M

mapX the sequence of the columns in original Mx.

mapY the sequence of the columns in original MY.

Ingca

Decompose the original data through LNGCA method.

Description

Implements the methods of linear non-Gaussian component analysis (LNGCA) and likelihood component analysis (when using a density, e.g., tilted Gaussian) from the LNGCA paper

Usage

```
lngca(
  xData,
  n.comp = NULL,
  Ux.list = NULL,
  whiten = c("sqrtprec", "eigenvec", "none"),
  maxit = 1000,
  eps = 1e-06,
  verbose = FALSE,
  restarts.pbyd = 0,
  restarts.dbyd = 0,
  distribution = c("JB", "tiltedgaussian", "logistic"),
  density = FALSE,
```

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```
out.all = FALSE,
orth.method = c("svd", "givens"),
df = 0,
stand = FALSE,
...
)
```

Arguments

xData the original dataset for decomposition, matrix of n x px.

n. comp the number of components to be estimated.

Ux.list list of user specified initial values for Ux. If null, will generate random orthog-

onal matrices. See restarts.pbyd and restarts.dbyd

whiten whitening method. Defaults to "svd" which uses the n left eigenvectors divided

by sqrt(px-1) by 'eigenvec'. Optionally uses the square root of the n x n "preci-

sion" matrix by 'sqrtprec'.

maxit max iteration, defalut = 1000

eps default = 1e-06

verbose default = FALSE

restarts.pbyd default = 0. Generates p x d random orthogonal matrices. Use a large number for

large datasets. Note: it is recommended that you run lngca twice with different seeds and compare the results, which should be similar when a sufficient number of restarts is used. In practice, stability with large datasets and a large number

of components can be challenging.

restarts.dbyd default = 0. These are d x d initial matrices padded with zeros, which results in

initializations from the principal subspace. Can speed up convergence but may

miss low variance non-Gaussian components.

distribution distribution methods with default to tilted Gaussian. "logistic" is similar to in-

fomax ICA, JB is capable of capture super and sub Gaussian distribution while being faster than tilted Gaussian. (tilted Gaussian tends to be most accurate, but

computationally much slower.)

density return the estimated tilted Gaussian density? default = FALSE

out.all default = FALSE

orth.method default = 'svd'. Method to generate random initial matrices. See [gen.inits()]

df default = 0, df of the spline used in fitting the non-parametric density. use df=8

or so for tilted gaussian. set df=0 for JB and logistic.

stand whether to standardize the data to have row and column means equal to 0 and

the row standard deviation equal to 1 (i.e., all variables on same scale). Often

used when combined with singR for data integration.

... other arguments to tiltedgaussian estimation

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Value

Function outputs a list including the following:

U matrix rx x n, part of the expression that Ax = Ux x Lx and Ax x Xc = Sx, where Lx is the whitener matrix.

loglik the value of log-likelihood in the lngca method.

S the variable loading matrix r x px, each row is a component, which can be used to measure nongaussianity

df egree of freedom.

distribution the method used for data decomposition.

whitener A symmetric whitening matrix n x n from dX, the same with whitener XA = est.sigmaXA%^% -0.5

M Mx Mtrix with n x rx.

nongaussianity the nongaussianity score for each component saved in S matrix.

```
#get simulation data
data(exampledata)
data=exampledata
# To get n.comp value, we can use NG_number function.
# use JB statistic as the measure of nongaussianity to run lngca with df=0
estX_JB = lngca(xData = data$dX, n.comp = 4,
whiten = 'sqrtprec', restarts.pbyd = 20, distribution='JB',df=0)
# use the tiltedgaussian distribution to run lngca with df=8. This takes a long time:
estX_tilt = lngca(xData = data$dX, n.comp = 4,
whiten = 'sqrtprec', restarts.pbyd = 20, distribution='tiltedgaussian',df=8)
# true non-gaussian component of Sx, include individual and joint components
trueSx = rbind(data$sjX,data$siX)
# use pmse to compare the difference of the two methods
pmse(S1 = t(trueSx),S2=t(estX_JB$S),standardize = TRUE)
pmse(S1 = t(trueSx),S2=t(estX_tilt$S),standardize = TRUE)
# the lngca using tiltedgaussian tends to be more accurate
# with smaller pmse value, but takes longer to run.
```

matchICA 13

matchICA match ICA

Description

match ICA

Usage

```
matchICA(S, template, M = NULL)
```

Arguments

S loading variable matrix template template for match M subject score matrix

Value

the match result

 NG_number

find the number of non-Gaussian components in the data.

Description

find the number of non-Gaussian components in the data.

Usage

```
NG_number(data, type = "S3")
```

Arguments

data original matrix with $n \times p$. type 'S1', 'S2' or 'S3'

Value

the number of non-Gaussian components in the data.

```
library(singR)
data("exampledata")
data=exampledata
NG_number(data$dX)
```

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orthogonalize

Orthogonalization of matrix

Description

Orthogonalization of matrix

Usage

```
orthogonalize(W)
```

Arguments

W

arbitrary matrix

Value

orthogonalized matrix

permmatRank_joint

Permutation test to get joint components ranks

Description

Permutation test to get joint components ranks

Usage

```
permmatRank_joint(matchedResults, nperms = 100)
```

Arguments

```
matchedResults results generated by angleMatchICA nperms the number of permutation
```

Value

a list of matrixes ## pvalues: pvalues for the matched columns don't have correlation. ## correpress: correlation value for original Mx with each random permutation of My. ## corrmatched: the correlation for each pair of matched columns.

permTestJointRank 15

permTestJointRank

Permutation test with Greedymatch

Description

Permutation test with Greedymatch

Usage

```
permTestJointRank(
  MatchedMx,
  MatchedMy,
  nperm = 1000,
  alpha = 0.01,
  multicore = 0
)
```

Arguments

MatchedMx matrix with nsubject x n.comp.X, comes from greedymatch matrix with nsubject 2 x n.comp.Y, comes from greedymatch

nperm $default \ value = 1000$ alpha $default \ value = 0.01$ multicore $default \ value = 0$

Value

a list of matrixes ## rj: joint component rank ## pvalues: pvalue for the components(columns) not matched ## fwer_alpha: quantile of corr permutation with 1- alpha

pmse

Permutation invariant mean squared error

Description

Permutation invariant mean squared error

Usage

```
pmse(M1 = NULL, M2 = NULL, S1 = NULL, S2 = NULL, standardize = FALSE)
```

signchange signchange

Arguments

M1	Subject score 1 matrix r x n.
M2	Subject score 2 matrix r x n.
S1	Loading 1 with matrix p x r.
S2	Loading 2 with matrix p x r.
standardize	whether to standardize

Value

permutation invariant mean squared error

Examples

```
#get simulation data
data(exampledata)

# use JB stat to compute with singR
output_JB=singR(dX=exampledata$dX,dY=exampledata$dY,
df=0,rho_extent="small",distribution="JB",individual=TRUE)

# use pmse to measure difference from the truth
pmse(M1 = t(output_JB$est.Mj),M2 = t(exampledata$mj),standardize = TRUE)
```

signchange

Sign change for S matrix to image

Description

Sign change for S matrix to image

Usage

```
signchange(S, M = NULL)
```

Arguments

```
 S \hspace{1cm} S, \, r \, x \, px.   M \hspace{1cm} Mx, \, n \, x \, r.
```

Value

a list of positive S and positive Mx.

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singR

SImultaneous Non-Gaussian Component analysis for data integration.

Description

This function combines all steps from the SING paper

Usage

```
singR(
 dΧ,
 dΥ,
 n.comp.X = NULL,
 n.comp.Y = NULL,
 df = 0,
  rho_extent = c("small", "medium", "large"),
 Cplus = TRUE,
  tol = 1e-10,
 stand = FALSE,
 distribution = "JB",
 maxiter = 1500,
 individual = FALSE,
 whiten = c("sqrtprec", "eigenvec", "none"),
 restarts.dbyd = 0,
  restarts.pbyd = 20
)
```

Arguments

C	XK	original dataset for decomposition, matrix of n x px.
C	Yk	original dataset for decomposition, matrix of n x py.
r	n.comp.X	the number of non-Gaussian components in dataset X . If null, will estimate the number using ICtest::FOBIasymp.
r	n.comp.Y	the number of non-Gaussian components in dataset Y. If null, will estimate the number using ICtest::FOBIasymp.
C	df	default value=0 when use JB, if df>0, estimates a density for the loadings using a tilted Gaussian (non-parametric density estimate).
r	ho_extent	Controls similarity of the scores in the two datasets. Numerical value and three options in character are acceptable. small, medium or large is defined from the JB statistic. Try "small" and see if the loadings are equal, then try others if needed. If numeric input, it will multiply the input by JBall to get the rho.
(Cplus	whether to use C code (faster) in curvilinear search.
t	tol	difference tolerance in curvilinear search.
S	stand	whether to use standardization, if true, it will make the column and row means to 0 and columns sd to 1. If false, it will only make the row means to 0.

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distribution "JB" or "tiltedgaussian"; "JB" is much faster. In SING, this refers to the "den-

sity" formed from the vector of loadings. "tiltedgaussian" with large df can

potentially model more complicated patterns.

maxiter the max iteration number for the curvilinear search.

individual whether to return the individual non-Gaussian components, default value = F.

whiten whitening method used in lngca. Defaults to "svd" which uses the n left eigen-

vectors divided by sqrt(px-1) by 'eigenvec'. Optionally uses the square root of

the n x n "precision" matrix by 'sqrtprec'.

restarts.dbyd default = 0. These are d x d initial matrices padded with zeros, which results in

initializations from the principal subspace. Can speed up convergence but may

miss low variance non-Gaussian components.

restarts.pbyd default = 20. Generates p x d random orthogonal matrices. Use a large number

for large datasets. Note: it is recommended that you run lngca twice with different seeds and compare the results, which should be similar when a sufficient number of restarts is used. In practice, stability with large datasets and a large

number of components can be challenging.

Value

Function outputs a list including the following:

Sjx variable loadings for joint NG components in dataset X with matrix rj x px.

Sjy variable loadings for joint NG components in dataset Y with matrix rj x py.

Six variable loadings for individual NG components in dataset X with matrix riX x px.

Siy variable loadings for individual NG components in dataset Y with matrix riX x py.

Mix scores of individual NG components in X with matrix n x riX.

Miy scores of individual NG components in Y with matrix n x riY.

est. Mjx Estimated subject scores for joint components in dataset X with matrix n x rj.

est. Mjy Estimated subject scores for joint components in dataset Y with matrix n x rj.

est.Mj Average of est.Mjx and est.Mjy as the subject scores for joint components in both datasets with matrix n x rj.

C_plus whether to use C version of curvilinear search.

rho_extent the weight of rho in search

df degree of freedom, = 0 when use JB, >0 when use tiltedgaussian.

```
#get simulation data
data(exampledata)

# use JB stat to compute with singR
output_JB=singR(dX=exampledata$dX,dY=exampledata$dY,
df=0,rho_extent="small",distribution="JB",individual=TRUE)
```

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```
# use tiltedgaussian distribution to compute with singR.
# tiltedgaussian may be more accurate but is considerably slower,
# and is not recommended for large datasets.
output_tilted=singR(dX=exampledata$dX,dY=exampledata$dY,
df=5,rho_extent="small",distribution="tiltedgaussian",individual=TRUE)
# use pmse to measure difference from the truth
pmse(M1 = t(output_JB$est.Mj),M2 = t(exampledata$mj),standardize = TRUE)

pmse(M1 = t(output_tilted$est.Mj),M2 = t(exampledata$mj),standardize = TRUE)
```

standard

Standardization with double centered and column scaling

Description

Standardization with double centered and column scaling

Usage

```
standard(data, dif.tol = 0.001, max.iter = 10)
```

Arguments

data input matrix with n x px.

dif. tol the value for the threshold of scaling

max.iter default value = 10

Value

standardized matrix with n x px.

```
spmwm = 3*matrix(rnorm(100000),nrow=100)+1
dim(spmwm)
apply(spmwm,1,mean) # we want these to be 0
apply(spmwm,2,mean) # we want these to be 0
apply(spmwm,2,sd) # we want each of these variances to be 1
spmwm_cp=standard(spmwm)
max(abs(apply(spmwm_cp,1,mean)))
max(abs(apply(spmwm_cp,2,mean)))
max(abs(apply(spmwm_cp,2,sd)-1))
```

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theta2W

Convert angle vector into orthodox matrix

Description

Convert angle vector into orthodox matrix

Usage

```
theta2W(theta)
```

Arguments

theta

vector of angles theta

Value

an orthodox matrix

tilted gaussian

tilted gaussian

Description

tiltedgaussian

Usage

```
tiltedgaussian(xData, df = 8, B = 100, ...)
```

Arguments

xData input data

df degree freedom

B default value=100

... ellipsis

vec2net 21

vec2net Create network matrices from vectorized lower diagonals vec2net transfer the matrix vectorized lower diagonals into net to show the

component image.

Description

Create network matrices from vectorized lower diagonals vec2net transfer the matrix vectorized lower diagonals into net to show the component image.

Usage

```
vec2net(invector, make.diag = 1)
```

Arguments

invector vectorized lower diagonals.

make.diag default value = 1.

Value

a net matrx

Examples

```
net = vec2net(1:10)
```

whitener

Whitening Function

Description

Whitening Function

Usage

```
whitener(X, n.comp = ncol(X), center.row = FALSE)
```

Arguments

X dataset p x n.

n.comp the number of components center.row whether center the row of data

Value

a whitener matrix

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%^%

Calculate the power of a square matrix

Description

returns a matrix composed of eigenvector x diag(eigenvalue ^ power) x eigenvector'

Usage

```
S %^% power
```

Arguments

S a square matrix power the times of power

Value

a matrix after power calculation that eigenvector x diag(eigenvalue ^ power) x eigenvector'

```
a <- matrix(1:9,3,3)
a %^% 2
```

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