

Package ‘MultiGroupO’

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Type Package

Title MultiGroup Method and Simulation Data Analysis

Version 0.4.0

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Description Two method new of multigroup and simulation of data.

The first technique called multigroup PCA (mgPCA) this multivariate exploration approach that has the idea of considering the structure of groups and / or different types of variables. On the other hand, the second multivariate technique called Multigroup Dimension-

ality Reduction (MDR) it is another multivariate exploration method that is based on projections. In addition, a method called Single Dimension Exploration (SDE) was incorporated for to analyze the exploration of the data. It could help us in a better way to observe the behavior of the multigroup data with certain variables of interest.

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Encoding UTF-8

Imports mvtnorm, rlist, expm, stats, ggplot2, gridExtra, cowplot,
plsgenomics, gplots, ggrepel, qgraph, mgm,lemon

Suggests knitr, rmarkdown

ByteCompile yes

VignetteBuilder knitr

RoxygenNote 7.2.3

NeedsCompilation no

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Contents

BIplot	2
fun.sim	3
mdr	4
mgpca	4
new.cov	5
pca	6
sde.method	7
Index	8

BIplot	<i>biplot methods</i>
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Description

biplot methods

Usage

```
BIplot(  
  variates,  
  loadings,  
  prop_expl_var,  
  comp = c(1, 2),  
  group = NULL,  
  rownamevar = T,  
  rownameload = T  
)
```

Arguments

- | | |
|---------------|--|
| variates | is the size of groups |
| loadings | is a vector of classes |
| prop_expl_var | data set |
| comp | component numeric |
| group | is a vector of groups |
| rownamevar | is a logical vector where TRUE is the label of the observations, if is FALSE, is index. |
| rownameload | is a logical vector where TRUE is the label of the vectors of loadings, if is FALSE, is index. |

Value

return an grafics .

Examples

```
library(datasets)
obj<-pca(datos=iris[, -5], grupos=iris[, 5], Plot=FALSE, center=TRUE, scale=TRUE)
BIplot(variates=obj$variates, loadings=obj$loadings,
        prop_expl_var=obj$prop_expl_var, comp=c(1, 2),
        group=factor(as.numeric(iris[, 5])), rownamevar=FALSE, rownameload=TRUE)
```

fun.sim	<i>Simulation function of quantitative multigroup data under a multivariate normal distribution</i>
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Description

Simulation function of quantitative multigroup data under a multivariate normal distribution

Usage

```
fun.sim(g, mean1, d, n.var, sds2, corr)
```

Arguments

g	An vector of the size of each group
mean1	An vector of the population means structure
d	distance d for the structure of population means
n.var	2x1 dimension vector whose first component is the number of random variables to simulate and the second component number of noise variables to simulate
sds2	An vector of the variances to simulate for each group noise variables
corr	An vector of the correlation to simulate for each group and noise variables

Value

return an grafics

Examples

```
fun.sim(g=c(20, 20), mean1=2, d=0, sds2=c(1, 1, 1), corr=c(0.5, 0.5, 0), n.var=c(50, 1))
```

mdr	<i>Performs a Multigroup Dimensionality Reduction (MDR) analysis in the given multigroup data matrix. Show MDR graphical output.</i>
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Description

Performs a Multigroup Dimensionality Reduction (MDR) analysis in the given multigroup data matrix. Show MDR graphical output.

Usage

```
mdr(group, data.x, c, Plot = T)
```

Arguments

group	is a vector of classes
data.x	quantitative data set
c	component numeric
Plot	grafics output of MDR

Value

return an grafics .

Examples

```
sim.list<-fun.sim(g=c(50,50,50),mean1=2,d=0,sds2=c(1,1,1),
corr=c(0.5,0.5,0.5,0),n.var=c(30,30))

mdr(group=as.factor(sim.list$grp),
data.x=sim.list$`lisx`,c=2)
```

mgpca	<i>Performs a Multigroup PCA analysis in the given multigroup data matrix. Show mgpca graphical output.</i>
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Description

Performs a Multigroup PCA analysis in the given multigroup data matrix. Show mgpca graphical output.

Usage

```
mgpca(
  mat.to.diag,
  mat.x,
  cls,
  Plot = TRUE,
  ncomp = 2,
  center = TRUE,
  scale = TRUE
)
```

Arguments

<code>mat.to.diag</code>	is a matrix with the data
<code>mat.x</code>	is a vector of classes
<code>cls</code>	group
<code>Plot</code>	grafics output of mgpca
<code>ncomp</code>	number of component
<code>center</code>	is a logical vector where TRUE is center (whether the variables should be shifted to be zero centered), if is FALSE, is original data.
<code>scale</code>	is a logical vector where TRUE is scale (indicating whether the variables should be scaled), if is FALSE, is original data.

Value

If `simplify == TRUE` class values. If `simplify == FALSE`, the result is a list of length `nsim` data.tables.

Examples

```
library(plsgenomics)
data(SRBCT)
mydata<-SRBCT$X
mydata<-mydata[1:50,1:5]
groups<-as.factor(SRBCT$Y)[1:50]
mat.to.diag1<-new.cov(x=mydata,cls=groups,A=diag(ncol(mydata)))
mgpca(mat.to.diag=mat.to.diag1,mat.x=as.matrix(mydata),
  cls=groups,Plot=TRUE,ncomp=2,center = TRUE,scale = TRUE)
```

new.cov

Function for the new covariance matrix in the multigroup PCA method

Description

Generates covariance matrix...

Usage

```
new.cov(x, cls, A)
```

Arguments

`x` is a matrix with the data
`cls` is a vector of classes
`A` is a symmetric and positive definite matrix associated to inner product respect to the base of its vectorial space.

Value

return an grafics.

Examples

```
library(plsgenomics)
data(SRBCT)
mydata<-SRBCT$X
mydata<-mydata[1:50,1:20]
groups<-as.factor(SRBCT$Y)[1:50]
new.cov(x=mydata,cls=groups,A=diag(ncol(mydata)))
```

pca	<i>Performs a principal components analysis in the given data matrix. Show PCA graphical output.</i>
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Description

Performs a principal components analysis in the given data matrix. Show PCA graphical output.

Usage

```
pca(datos, grupos, Plot = TRUE, center = TRUE, scale = TRUE)
```

Arguments

`datos` is a matrix with the data
`grupos` is a vector of classes
`Plot` vector logic for grafic
`center` data set center by columns
`scale` data set scaled by columns

Value

return an grafics.

Examples

```
library(plsgenomics)
data(SRBCT)
mydata<-SRBCT$X
mydata<-mydata[1:30,1:20]
groups<-as.factor(SRBCT$Y)[1:30]
pca(datos=mydata, grupos=groups, Plot=TRUE, center=TRUE, scale=TRUE)
```

sde.method	<i>Performs a Single Dimension Exploration (SDE) analysis in the given multigroup data matrix. Show SDE graphical output.</i>
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Description

Performs a Single Dimension Exploration (SDE) analysis in the given multigroup data matrix. Show SDE graphical output.

Usage

```
sde.method(mydata, groups, plt = FALSE)
```

Arguments

mydata	data set
groups	is a vector of classes
plt	grafics

Value

return an grafics .

Examples

```
sim.list2<-fun.sim(g=c(20,50,10),mean1=0.5,d=0,sds2=c(1,1,1,1),corr=c(0.1,0.5,0.5,0),
n.var=c(20,20))
datos2 <- as.data.frame(sim.list2$x)
datos2<-subset(datos2,select=-grp)
grupos <- sim.list2$grp
grupos<-factor(grupos,labels=c(1,2,3))
sde.method(mydata=datos2,groups=grupos,plt=FALSE)
```

Index

BIplot, [2](#)

fun.sim, [3](#)

mdr, [4](#)

mgpca, [4](#)

new.cov, [5](#)

pca, [6](#)

sde.method, [7](#)