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Description Tools of Bayesian analysis framework using the method suggested by Berger (1985) <doi:10.1007/978-1-4757-4286-2> for multivariate normal (MVN) distribution and multivariate normal mixture (MixMVN) distribution: a) calculating Bayesian posteriori of (Mix)MVN distribution; b) generating random vectors of (Mix)MVN distribution; c) Markov chain Monte Carlo (MCMC) for (Mix)MVN distribution.

Imports mvtnorm, plyr, stats

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MVNBayesian-package Bayesian Analysis Framework for MVN (Mixture) Distribution

Description

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Tools of Bayesian analysis framework using the method suggested by Berger (1985) <doi:10.1007/978-1-4757-4286-2> for multivariate normal (MVN) distribution and multivariate normal mixture (MixMVN) distribution: a) calculating Bayesian posteriori of (Mix)MVN distribution; b) generating random vectors of (Mix)MVN distribution; c) Markov chain Monte Carlo (MCMC) for (Mix)MVN distribution.

Details

This package is aimed to build a easy approach for MVN (mixture) distribution in Bayesian analysis framework. Bayesian posteriori MVN (mixture) distribution can be calculated in conditions of given priori MVN (mixture) informations. The conjugated property of MVN distribution makes it effective in parameter estimation using Bayesian iterator. Joint and marginal probability densities of a certain MVN (mixture) can be achieved through random vector generator, using Gibbs sampling. Conditional probability densities from a certain MVN (mixture) can be simulated using MCMC method.

Author(s)

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References

"Statistical Inference" by George Casella. Roger L. Berger;

"Statistical Decision Theory and Bayesian Analysis" by James O. Berger;

"Matrix Computation" by Gee H. Golub. Charles F. Van Loan;

"Bayesian Statistics" by WEI Laisheng;

"Machine Learning" by NAKAGAWA Hiroshi.

Ascending_Num

See Also

stats, mvtnorm

Examples

```
library(Rfast)
library(mvtnorm)
library(plyr)
head(dataset1)
BP <- MVN_BayesianPosteriori(dataset1)
BP
BP_Gibbs <- MVN_GibbsSampler(5000, BP)
colMeans(BP_Gibbs)
colrange(BP_Gibbs)
result <- MVN_MCMC(BP, 5000, c(1), c(77.03))
result$Accept</pre>
```

Ascending_Num Renumbering vector by elemental frequency

Description

Renumbering vector by elemental frequency in ascending order.

Usage

Tidy vector by elemental frequency: Ascending_Num(data)

Arguments

data An 1d-vector.

Value

return a renumbered vector by elemental frequency. Factors will be positive integers arrayed in ascending order.

Examples

library(plyr)
x <- c(1,2,2,2,2,2,2,2,3,3,3,1,3,3,3)
x
Ascending_Num(x)</pre>

dataset1

Description

Dataset built for MVN mixture test, which contains 3 variables and 25 observations.

Usage

data("dataset1")

Format

A data frame with 25 observations on 3 independent variables, named as fac1, fac2 and fac3.

fac1 The 1st factor.

fac2 The 2nd factor.

fac3 The 3rd factor.

Examples

dataset1

dataset2

Dataset for MVN mixture test

Description

Dataset built for MVN mixture test, which contains 4 variables (the first 4 columns), clustering (the last column) and 96 observations.

Usage

data("dataset2")

Format

A data frame with 96 pseudo-observations generated by random number generator. All observations come from 3 different centers which have been marked in the last column "species". More specifically, data of species=1 comes from the center (1,1,1,1); data of species=2 comes from the center (2,2,2,0); data of species=3 comes from the center (1,0,2,2).

dimen1 the 1st variable

- dimen2 the 2nd variable
- dimen3 the 3rd variable
- dimen4 the 4th variable
- species clustering label

MatrixAlternative

Examples

dataset2

MatrixAlternative Interchanging specified rows and columns

Description

Interchange all elements between two specified rows and columns in a matrix.

Usage

A matrix-like data
MatrixAlternative(data, sub, rep)

Arguments

data	A matrix to be processed.
sub	A positive integer. The first selected dimension.
rep	A positive integer. The second selected dimension. Default value is 1

Value

return a matrix with interchanged rows and columns in two specified dimensions.

Examples

```
library(plyr)
M <- matrix(1:9,3,3,1)
M
MatrixAlternative(M, 2)</pre>
```

MixMVN_BayesianPosterior

MixMVN_BayesianPosterior

Description

Given a design matrix (data) including sufficient samplings from different multivariate normal distribution, export the parameters of Bayesian posterior multivariate normal mixture distribution. Parameters contains mixture probability, mean vector and covariance matrix, for each cluster.

Usage

```
# paramtric columns-only as input data:
# data <- dataset2[,1:4]
# Specify species to get parameters of mixture MVN:
MixMVN_BayesianPosterior(data, species, idx)
```

Arguments

data	Design matrix: data.frame or matrix-like data sampling from different multivari- ate normal distribution, dim(data)[1] should be the number of observations and dim(data)[2] should be the number of variables or parameters.
species	Number of clusters: number of clusters for import data. It will be only called once by the next argument idx through kmeans clustering algrithm in this func- tion. Default value 1 means no clustering is used if the idx is also the default. In that case, this function will execute identical process as MVN_BayesianPosterior()
idx	port for clustering result: an vector which should have the identical dimension as the number of samplings of data. Default value is generated by kmeans algrithm.

Value

return a matrix-like result with contains all parameters of Bayesian posterior multivariate normal mixture distribution: All clusters are arrayed in rows and mixture probability, posterior mean and posterior covariance for each cluster are arrayed in columns.

Author(s)

ZHANG Chen

See Also

kmeans, MVN_BayesianPosterior

Examples

library(plyr)

```
# Design matrix should only contain columns of variables
# Export will be a matrix-like data
# Using kmeans (default) clustering algrithm
data_dim <- dataset2[,1:4]
result <- MixMVN_BayesianPosterior(data=data_dim, species=3)
result
# class(result)
# Get the parameters of the 1st cluster:
result[1,]
# class(result[1,])
```

Get the mixture probability of cluster2:

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```
result[2,1][[1]]
# class(result[2,1][[1]])
# class(result[2,1]$probability)
# class(result[2,1])
# Get the mean vector of cluster1:
result[1,2][[1]]
# class(result[1,2][[1]])
# class(result[1,2])
# Get the covariance matrix of cluster3:
result[3,3][[1]]
# class(result[3,3][[1]])
# class(result[3,3][1]])
# class(result[3,3][1]])
# class(result[3,3])
```

MixMVN_BayesianPosteriori Calculate Bayesian posteriori MVN mixture distribution

Description

The function to export the mixture probabilities, the mean vectors and covariance matrices of Bayesian posteriori MVN mixture distribution in the basis of given priori information (priori MVN mixture) and observation data (a design matrix containing all variables).

Usage

```
# paramtric columns-only as input data:
# data <- dataset2[,1:4]</pre>
```

Specify species to get parameters of MVN mixture model: MixMVN_BayesianPosteriori(data, species, idx)

Arguments

data	A data.frame or matrix-like data: observations should be arrayed in rows while variables should be arrayed in columns.
species	A positive integer. The number of clusters for import data. It will be only called once by the next argument idx through kmeans clustering algrithm in this function. Default value is 1, which means no clustering algrithm is used.
idx	A vector-like data to import for accepting clustering result. Default value is generated by kmeans clustering. Notice the length of idx should be the same as observation numbers of data (rows).

Value

return a matrix-like result containing all parameters of Bayesian posteriori MVN mixture distribution: Clusters are arrayed in rows, while the mixture probabilities, posteriori mean vectors and posteriori covariance matrices are arrayed in columns.

See Also

kmeans, MVN_BayesianPosteriori

Examples

library(plyr)

```
# Design matrix should only contain columns of variables
# Export will be a matrix-like data
# Using kmeans (default) clustering algrithm
data_dim <- dataset2[,1:4]</pre>
result <- MixMVN_BayesianPosteriori(data=data_dim, species=3)</pre>
result
# Get the parameters of the cluster1:
result[1,]
# Get the mixture probability of cluster2:
# (Attention to the difference between
# result[2,1][[1]] and result[2,1])
result[2,1][[1]]
# Get the mean vector of cluster1:
result[1,2][[1]]
# Get the covariance matrix of cluster3:
result[3,3][[1]]
```

MixMVN_GibbsSampler Gibbs sampler for MVN mixture distribution

Description

Generating random vectors on the basis of a given MVN mixture distribution, through Gibbs sampling algorithm or matrix factorization.

Usage

```
# Bayesian posteriori MVN mixture model as input data:
# data <- MixMVN_BayesianPosteriori(dataset2[,1:4], species=3)</pre>
```

Generate random vectors based on Bayesian posteriori MVN mixture: MixMVN_GibbsSampler(n, data, random_method = c("Gibbs", "Fast"), reject_rate=0, ...)

Arguments

n	A positive integer. The numbers of random vectors to be generated.
data	A matrix-like data which contains the mixture probability, mean vector and co- variance matrix for each cluster in each row.
random_method	The method to generate random vectors. Options are "Gibbs": Gibbs sampling for MVN mixture model; and "Fast": call rmvnorm() to generate random vectors based on matrix factorization.
reject_rate	A numeric value which will be efficient if the random_method is "Gibbs": De- termine the discarded items in burn-in period by ratio. Default value is 0. For details see MVN_GibbsSampler.
	Other arguments to control the process in Gibbs sampling if the random_method is "Gibbs".

Details

It is recommanded using the random method of "Fast" due to the high efficiency. The time complexity of "Gibbs" method is O(k*n) where the k means dimensionality of MVN mixture model and n means generated numbers of random vectors; while that of the "Fast" method is only O(n), without considering the effect of burn-in period. this discrepancy will be even further significant when we use MCMC methods to do some further analysis in which random vectors will be generated every time when we set conditions.

Value

return a series random vectors in the basis of given MVN mixture distribution.

See Also

Ascending_Num, MixMVN_BayesianPosteriori, MVN_BayesianPosteriori

Examples

```
library(plyr)
library(mvtnorm)
library(stats)
# Use dataset2 for demonstration. Get parameters of Bayesian
# posteriori multivariate normal mixture distribution
head(dataset2)
dataset2_par <- dataset2[,1:4] # only parameter columns are premitted
MixBPos <- MixMVN_BayesianPosteriori(dataset2_par, species=3)
MixBPos
# Generate random vectors using Gibbs sampling:
MixBPos_Gibbs <- MixMVN_GibbsSampler(5000, MixBPos, random_method = "Gibbs")
head(MixBPos_Gibbs)</pre>
```

```
# Compared generation speed of "Gibbs" to that of "Fast"
MixBPos_Fast <- MixMVN_GibbsSampler(5000, MixBPos, random_method = "Fast")</pre>
```

```
head(MixBPos_Fast)
# Visulization by clusters:
library(rgl)
dimen1 <- MixBPos_Gibbs[,1]
dimen2 <- MixBPos_Gibbs[,2]
dimen3 <- MixBPos_Gibbs[,3]
dimen4 <- MixBPos_Gibbs[,4]
plot3d(x=dimen1, y=dimen2, z=dimen3, col=MixBPos_Gibbs[,5], size=2)</pre>
```

MixMVN_MCMC

MCMC simulation for MVN mixture distribution

Description

Function to get a MCMC simulation results based on the imported MVN mixture distribution. It is commonly used for inquiring the specified conditional probability of MVN mixture distribution calculated through Bayesian posteriori.

Usage

```
# Bayesian posteriori mix MVN as input data:
# data <- MixMVN_BayesianPosteriori(dataset2[,1:4], 3)</pre>
```

```
# run MCMC simulation based on Bayesian posteriori mix MVN:
MixMVN_MCMC(data, steps, pars, values, tol, random_method, ...)
```

Arguments

data	A matrix-like data containing the mixture probability, mean vector and covari- ance matrix for each cluster in each row.
steps	A positive integer. The numbers of random vectors to be generated for MCMC step.
pars	A integer vector to declare fixed dimension(s). For example if the desired dimensions are 1st=7 and 3rd=10, set this argument as $c(1,3)$.
values	A numeric vector to assign value(s) to declared dimension(s). For example if the desired dimensions are $1st=7$ and $3rd=10$, set this argument as $c(7,10)$.
tol	Tolerance. A numeric value to control the generated vectors to be accepted or rejected. Criterion uses Euclidean distance in declared dimension(s). Default value is 0.3.
random_method	The method to generate random vectors. Options are "Gibbs": Gibbs sampling for MVN mixture model; and "Fast": call rmvnorm() to generate random vectors based on matrix factorization. Default option is "Fast".
	Other arguments to control the process in Gibbs sampling if the random_method is "Gibbs".

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MixMVN_MCMC

Value

return a list which contains:

AcceptRate	Acceptance of declared conditions of MCMC
MCMCdata	All generated random vectors in MCMC step based on MVN mixture distribution
Accept	Subset of accepted sampling in MCMCdata
Reject	Subset of rejected sampling in MCMCdata

See Also

MixMVN_BayesianPosteriori, MixMVN_GibbsSampler, MVN_GibbsSampler, MVN_FConditional

Examples

```
library(plyr)
library(mvtnorm)
library(stats)
# dataset2 has 4 parameters: dimen1, dimen2, dimen3 and dimen4:
head(dataset2)
dataset2_dim <- dataset2[,1:4] # extract parametric columns</pre>
# Get posteriori parameters of dataset2 using kmeans 3 clustering:
MixBPos <- MixMVN_BayesianPosteriori(dataset2_dim, 3)</pre>
# If we want to know when dimen1=1, which clusters are accepted, run:
MixBPos_MCMC <- MixMVN_MCMC(MixBPos, steps=5000, pars=c(1), values=c(1), tol=0.3)</pre>
MixBPos_MCMC$AcceptRate
result <- MixBPos_MCMC$MCMCdata</pre>
head(result)
# count accepted samples by clustering:
count(result[which(result[,7]==1),5])
library(rgl)
# Visualization using plot3d() if necessary:
# Clustering result in the rest 3 dimensions:
plot3d(result[,2], result[,3], z=result[,4], col=result[,5], size=2)
# Acceptance rejection visualization:
plot3d(result[,2], result[,3], z=result[,4], col=result[,7]+1, size=2)
```

MVN_BayesianIterator Parameter estimation using Bayesian iteration

Description

Function to execute parameter estimation for MVN distribution, under Bayesian analysis frame-work.

Usage

```
# Get parameters of Bayesian posteriori MVN:
MVN_BayesianIterator(data, pri_mean=colMeans(data), Gibbs_nums=5000,
pseudo_nums=dim(data)[1], threshold=1e-04, iteration=100, ...)
```

Arguments

data	A data.frame or matrix-like data: obervations should be arrayed in rows while variables should be arrayed in columns.	
pri_mean	A numeric vector to assign priori mean for MVN. Default value applies colMeans() to data.	
Gibbs_nums	A positive integer. The numbers of random vectors to be generated for each iteration step. Defult value is 5000.	
pseudo_nums	A positive integer. The argument to determine numbers of generated vectors used for each iteration step. Default value keeps the same scale as input data. Notice that a too small value can result in singular matrix.	
threshold	A numeric value to control stoping the iteration loop. Default value used 0.0001. While the Euclidean distance of mean vectors between pseudo-data (the last pseudo_nums items) and Bayesian posteriori is less than threshold, iteration stops.	
iteration	A positive integer. Argument to assign the maximum steps for iteration. Default value is 100 after which the iteration loop will compulsively exit.	
	Other arguments to control the process in Gibbs sampling.	

Details

Because that MVN distribution possess conjugated property in Bayesian analysis framework, the convergence of Bayesian iterator for MVN distribution can be ensured, accoumpanied with the shrink of 2nd-norm of Bayesian posteriori covariance matrix. But pay attention to the fact that pseudo-data leads to the randomness, the argument pseudo_nums should be set carefully.

Value

return a double level list containing Bayesian posteriori after iteration process:

mean	Bayesian posteriori mean vector
var	Bayesian posteriori covariance matrix

Note

If the parameter values are the only interested thing we concerned, this iterator makes sense. Since it can significantly help us decrease the scale of covariance matrix, to obtain a more reliable estimation for the parameters. However, in more cases, some correlationships of a certain group of pamameters are more valuable, which are usually clued by the covariance matrix.

See Also

MVN_BayesianPosteriori, MVN_GibbsSampler, MVN_FConditional, MatrixAlternative

Examples

library(mvtnorm)

```
# Bayesian posteriori before iteration using dataset1 as example,
# c(80, 16, 3) as priori mean:
# View 2-norm of covariance matrix of Bayesian posteriori:
BPos_init <- MVN_BayesianPosteriori(dataset1, c(80,16,3))
BPos_init
norm(as.matrix(BPos_init$var), type = "2")
# Bayesian posteriori after iteration using c(80,16,3) as priori
# Using 30 last samples generated by GibbsSampler for each step:
BPos_fina1 <- MVN_BayesianIterator(dataset1, c(80,16,3), 5000, 30)
BPos_fina1
```

norm(as.matrix(BPos_fina1\$var), type = "2")

Too small pseudo_nums setting can results in singular system, try: MVN_BayesianIterator(dataset1, pseudo_nums=3)

MVN_BayesianPosterior MVN_BayesianPosterior

Description

Given a design matrix (data) and priori information, export the mean vector and covariance matrix of Bayesian posterior multivariate normal distribution.

Usage

```
# Given the data as design matrix, priori mean vector and priori covariance
# matrix, this function will export a list which contains mean ($mean) and
# covariance ($var) of Bayesian posterior multivariate normal distribution.
MVN_BayesianPosterior(data, pri_mean, pri_var)
# defualt pri_mean uses colMenas()
```

defualt pri_var uses unit matrix

Arguments

data	Design matrix: data.frame or matrix-like data, dim(data)[1] should be the number of observations and dim(data)[2] should be the number of variables or parameters.
pri_mean	priori mean: necessary vector which should be of the identical dimensions of data (length(pri_mean) == dim(data)[2]). Users can set according to their own prior knowledge.
pri_var	prior covariance matrix: a real symmetric matrix by definition; the default value is an unit matrix with the same dimension of priori mean vector.

Details

Although this function is very simple, the observation data should be diagnosed firstly. it is strongly recommanded that researchers and developers should have some prior knowledge of ill-conditioned system before using this function. Simply, ill-conditioned system, or singular matrix, is caused by a) insufficient data or b) almostly linear dependency of two certain parameters, which two can result in a too small eigenvalue then cause a ill-conditioned (singular) system. Therefore users should make sure the data contains enough observations and the degree of freedom is strictly equal to the number of parameters.

Value

return a list of:

mean	mean vector of Bayesian posterior
var	covariance of Bayesian posterior

Author(s)

ZHANG Chen

Examples

```
# Demo using dataset1:
head(dataset1)
BPos <- MVN_BayesianPosterior(dataset1, c(80,16,3))
BPos$mean
BPos$var
```

```
# Singular system caused by insufficient data
eigen(var(dataset1[1:3,]))$values
rcond(var(dataset1[1:3,]))
eigen(var(dataset1[1:6,]))$values
rcond(var(dataset1[1:6,]))
```

```
# Singular system caused by improper degree of freedom
K <- cbind(dataset1, dataset1[,3]*(-2)+3)
eigen(var(K[,2:4]))$values
rcond(var(K[,2:4]))</pre>
```

MVN_BayesianPosteriori

Calculate Bayesian posteriori MVN distribution

Description

The function to export the mean vector and covariance matrix of Bayesian posteriori MVN distribution in the basis of given priori information (priori MVN) and observation data (a design matrix containing all variables).

Usage

Given the data as design matrix, priori mean vector and priori covariance # matrix, this function will export a list which contains mean (\$mean) and # covariance (\$var) of Bayesian posteriori multivariate normal distribution.

MVN_BayesianPosteriori(data, pri_mean, pri_var)

Arguments

data	A data.frame or matrix-like data: obervations should be arrayed in rows while variables should be arrayed in columns.
pri_mean	A numeric vector to assign priori mean for MVN. Default value applies colMeans() to data.
pri_var	A matrix-like parameter to assign priori covariance matrix. Default value uses unit matrix.

Value

return a double level list containing:

mean	mean vector of Bayesian posteriori MVN distribution
var	covariance of Bayesian posteriori MVN distribution

Note

It is strongly recommanded that users should have some prior knowledge of ill-conditioned system before using this function. Simply, ill-conditioned system, or singular matrix, is caused by a) insufficient data or b) almostly linear dependency of two certain parameters, which two can result in a excessively small eigenvalue then cause a ill-conditioned (singular) system. Therefore users must diagnose their data firstly to confirm the fact that the it contains enough observations, and the degree of freedom is strictly equal to the number of parameters as well. Additionally, for the argument pri_var, a real symmetric matrix is desired by definition.

Examples

```
# Demo using dataset1:
head(dataset1)
BPos <- MVN_BayesianPosteriori(dataset1, c(80,16,3))
BPos$mean
BPos$var
# Singular system caused by insufficient data
eigen(var(dataset1[1:3,]))$values
rcond(var(dataset1[1:6,]))$values
rcond(var(dataset1[1:6,]))
# Singular system caused by improper degree of freedom
K <- cbind(dataset1, dataset1[,3]*(-2)+3)
eigen(var(K[,2:4]))$values
rcond(var(K[,2:4]))
```

MVN_FConditional Calculate full conditional normal ditribution of MVN

Description

Function to export parameters of full conditional normal distribution in basis of given MVN distribution, the undecided dimension, as well as all values in the rest dimensions.

Usage

```
# Bayesian posteriori as input data:
# data <- MVN_BayesianPosteriori(dataset1, c(80,16,3))</pre>
```

inquire parameters of full-conditional distribution based on Bayesian posteriori: MVN_FConditional(data, variable, z)

Arguments

data	A double level list containing all parameters of MVN distribution: mean vector (data\$mean) and covariance matrix (data\$var).
variable	A integer to specify the undecided dimension.
Z	A nd-vector to assign conditions (n = dimensions of given MVN distribution). It should be noted that the value in dimension specified by variable doesn't participate in the calculation.

Details

It can be proved that any full conditional distribution from a given MVN will degenerate to an 1d-normal distribution.

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MVN_GibbsSampler

Value

return a double level list containing the following parameters of full conditional normal distributions of given MVN in specified dimension:

mean	a numberic mean of a normal distribution
var	a numberic variance of a normal distribution

See Also

MVN_BayesianPosteriori, MatrixAlternative

Examples

```
head(dataset1)
BPos <- MVN_BayesianPosteriori(dataset1, c(80,16,3))
BPos # Bayesian Posteriori
result <- MVN_FConditional(BPos, variable = 1, z=c(75, 13, 4))
result$mean
class(result$mean)
result$var
class(result$var)
# compare the following results:</pre>
```

```
MVN_FConditional(BPos, variable = 2, z=c(75, 13, 4))
MVN_FConditional(BPos, variable = 2, z=c(75, 88, 4))
MVN_FConditional(BPos, variable = 1, z=c(75, 88, 4))
```

MVN_GibbsSampler Gibbs sampler for MVN distribution

Description

Generating random vectors on the basis of a given MVN distribution, through Gibbs sampling algorithm.

Usage

```
# Bayesian posteriori as data
# data <- MVN_BayesianPosteriori(dataset1)
# Using Gibbs sampler to generate random vectors based on Bayesian posteriori:
MVN_GibbsSampler(n, data, initial, reject_rate, burn)</pre>
```

Arguments

n	A positive integer. The numbers of random vectors to be generated.
data	A double level list which contains the mean vector (data $mean$) and the covariance matrix (data var) of a given MVN distribution.
initial	Initial vector where Markov chain starts. Default value use a random vector generated by rmvnorm().
reject_rate	A numeric to control burn-in period by ratio. Default value is 0.2, namely the first 20% generated vectors will be rejected. If this arg was customized, the next arg burn should maintain the default value.
burn	A numeric to control burn-in period by numbers. If this arg was customized, final result will be generated by this manner in which it will drop the first n numbers (n=burn).

Details

There're also some literatures suggest using the mean or mode of priori as initial vector. Users can customize this setting according to their own needs.

Value

return a series random vectors in the basis of given MVN distribution.

See Also

MVN_FConditional, MatrixAlternative

Examples

library(mvtnorm)

```
# Get parameters of Bayesian posteriori multivariate normal distribution
BPos <- MVN_BayesianPosteriori(dataset1)
BPos
```

```
# Using previous result (BPos) to generate random vectors through Gibbs
# sampling: 7000 observations, start from c(1,1,2), use 0.3 burning rate
BPos_Gibbs <- MVN_GibbsSampler(7000, BPos, initial=c(1,1,2), 0.3)
tail(BPos_Gibbs)
```

```
# Check for convergence of Markov chain
BPos$mean
colMeans(BPos_Gibbs)
BPos$var
var(BPos_Gibbs)
```

```
# 3d Visulization:
library(rgl)
fac1 <- BPos_Gibbs[,1]
fac2 <- BPos_Gibbs[,2]</pre>
```

MVN_MCMC

```
fac3 <- BPos_Gibbs[,3]
plot3d(x=fac1, y=fac2, z=fac3, col="red", size=2)</pre>
```

MVN_MCMC

MCMC simulation for MVN distribution

Description

Function to get a MCMC simulation results based on the imported MVN distribution. It is commonly used for inquiring the specified conditional probability of MVN distribution calculated through Bayesian posteriori.

Usage

```
# Bayesian posteriori as input data
# data <- MVN_BayesianPosteriori(dataset1, pri_mean=c(80,16,3))</pre>
```

```
# run MCMC simulation using Bayesian posteriori:
MVN_MCMC(data, steps, pars, values, tol, ...)
```

Arguments

data	A double level list which contains the mean vector (data $mean$) and the covariance matrix (data var) of a given MVN distribution.
steps	A positive integer. The numbers of random vectors to be generated for MCMC step.
pars	A integer vector to declare fixed dimension(s). For example if the desired dimensions are $1st=7$ and $3rd=10$, set this argument as $c(1,3)$.
values	A numeric vector to assign value(s) to declared dimension(s). For example if the desired dimensions are $1st=7$ and $3rd=10$, set this argument as $c(7,10)$.
tol	Tolerance. A numeric value to control the generated vectors to be accepted or rejected. Criterion uses Euclidean distance in declared dimension(s). Default value is 0.3.
	Other arguments to control the process in Gibbs sampling.

Value

return a list which contains:

AcceptRate	Acceptance of declared conditions of MCMC
MCMCdata	All generated random vectors in MCMC step based on MVN distribution
Accept	Subset of accepted sampling in MCMCdata
Reject	Subset of rejected sampling in MCMCdata

See Also

MVN_GibbsSampler, MVN_FConditional

Examples

```
library(mvtnorm)
library(plyr)
```

dataset1 has three parameters: fac1, fac2 and fac3: head(dataset1)

```
# Get posteriori parameters of dataset1 using prior of c(80,16,3):
BPos <- MVN_BayesianPosteriori(dataset1, pri_mean=c(80,16,3))</pre>
```

```
# If we want to know when fac1=78, how fac2 responses to fac3, run:
BPos_MCMC <- MVN_MCMC(BPos, steps=8000, pars=c(1), values=c(78), tol=0.3)
MCMC <- BPos_MCMC$MCMCdata
head(MCMC)
```

```
# Visualization using plot3d() if necessary:
library(rgl)
plot3d(MCMC[,1], MCMC[,2], z=MCMC[,3], col=MCMC[,5]+1, size=2)
```

```
# Visualization: 2d scatter plot
MCMC_2d <- BPos_MCMC$Accept
head(MCMC_2d)
plot(MCMC_2d[,3], MCMC_2d[,2], pch=20, col="red", xlab = "fac3", ylab = "fac2")
```

```
# Compared to the following scatter plot when fac1 is not fixed:
plot(BPos_MCMC$MCMCdata[,3], BPos_MCMC$MCMCdata[,2], pch=20, col="red", xlab = "fac3",
ylab = "fac2")
```

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