

Package ‘BANFF’

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Type Package

Title Bayesian Network Feature Finder

Version 1.1

Description Provides a full package of posterior inference, model comparison, and graphical illustration of model fitting. A parallel computing algorithm for the Markov chain Monte Carlo (MCMC) based posterior inference and an Expectation-Maximization (EM) based algorithm for posterior approximation are developed, both of which greatly reduce the computational time for model inference.

Depends R (>= 3.0.3), foreach, doParallel

License GPL-2

Imports DPpackage, igraph, mclust, pscl, tmvtnorm, network, coda, graphics, methods, stats, utils

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DPM.HODC	<i>Hierarchical Ordered Density Clustering (HODC) for Dirichlet Process Mixture Model Fitting</i>
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Description

This function implements the HODC algorithm for posterior density samples by a Dirichlet process mixture (DPM) of normals model which is implemented by function "DPdensity" in R package **DPpackages**.

Usage

```
DPM.HODC(v, pvalue,
DPM.mcmc=list(nburn=2000,nsave=1,
nskip=0,ndisplay=10),
DPM.prior=list(a0=2,b0=1,m2=rep(0,1),
s2=diag(100000,1),
psiinv2=solve(diag(0.5,1)),
nu1=4,nu2=4,tau1=1,tau2=100))
```

Arguments

pvalue	a vector of p-values obtained from large scale statistical hypothesis testing
v	the number of posterior sample saved
DPM.mcmc	a list giving the MCMC parameters for DPM fitting; see the argument mcmc of function DPdensity() in DPpackage for details; the default setting is DPM.mcmc=list(nburn=2000,nsave=1,nskip=0,ndisplay=10)
DPM.prior	a list giving the prior information; see the argument prior of function DPdensity() in DPpackage for details; the default setting is prior2

Details

This function calls DPdensity to estimate the marginal density of the testing statistics \mathbf{r} , converted from p-values, using a mixture of normal densities without incorporating the network information. Furthermore, it implements the HODC algorithm to classify density components into two clusters. We refer to them as the unimportant cluster and the important cluster, where the important cluster has a larger mean than the unimportant cluster.

Value

a list of density clustering results by the HODC algorithm

mean a list containing posterior samples for the mean of unimportant and important clusters

mu0 a vector of length "v" containing posterior samples for the mean of the unimportant cluster

mu1 a vector of length "v" containing posterior samples for the mean of the important cluster

variance a list containing posterior samples of the variance of the unimportant and the important clusters

var0 a vector of length "v" containing posterior samples for the variance of the unimportant cluster

var1 a vector of length "v" containing posterior samples for the variance of the important cluster

probability a list containing the probabilities of unimportant and important clusters

pro0 a vector of length "v" containing posterior samples for the probability of the unimportant cluster

pro1 a vector of length "v" containing posterior samples for the probability of the important cluster

classification a binary (0/1) matrix of dimension "v" by length(pvalue) containing posterior samples for two cluster classification results

Author(s)

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Department of Biostatistics and Bioinformatics, Emory University

References

Yize Zhao, Jian Kang, Tianwei Yu (2014) A Bayesian nonparametric model for selecting gene and gene sub-network, *Annals of Applied Statistics*, in press.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Examples

```
###random make the density
rstat=c(rnorm(50,mean=1),rnorm(50,mean=2),rnorm(100,mean=4)
,rnorm(100,mean=8))
###transformed into pvalue
pvalue=pnorm(-rstat)
DPMHODC=DPM.HODC(v=5,pvalue)
```

EM.HODC

Hierarchical Ordered Density Clustering (HODC) for Finite Mixture Model Fitting

Description

This function implements the HODC algorithm for density estimates by a finite mixture of normals model which is implemented by function `Mclust` in R package **mclust**.

Usage

```
EM.HODC(pvalue)
```

Arguments

`pvalue` a vector of p-values obtained from large scale statistical hypothesis testing

Details

This function calls `Mclust` to estimate the marginal density of the testing statistics \mathbf{r} , converted from p-values, using a mixture of normal densities without incorporating the network information. Furthermore, it implements the HODC algorithm to classify density components into two clusters. We refer to them as the unimportant cluster and the important cluster, where the important cluster has a larger mean than the unimportant cluster.

Value

a list of density clustering results by the HODC algorithm

mean mean estimates of unimportant and important clusters

variance variance estimates of unimportant and important clusters

pro probability estimates of unimportant and important clusters

classification classifications configurations

Author(s)

Zhou Lan, Jian Kang, Tianwei Yu and Yize Zhao

References

Yize Zhao, Jian Kang, Tianwei Yu (2014) A Bayesian non parametric model for selecting gene and gene sub network, *Annals of Applied Statistics*, in press.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Examples

```
rstat=c(rnorm(50,mean=1),rnorm(50,mean=2),rnorm(100,mean=4),rnorm(100,mean=8))
pvalue=pnorm(-rstat)
EMHODC=EM.HODC(pvalue)
```

```
Grid.Adjmatrix.Transfer
```

Connector for Transferring Grid Coordinates into Adjacency Matrix

Description

This function is a supplementary function for transferring grid coordinates into adjacency matrix. The criteria for defining connected and disconnected: for any two vectors, if euclidean distance is less than a certain value, we consider these two vectors as connected, otherwise, disconnected. The purpose of this function is to quickly transform image data, such as brain image data, into adjacency matrix.

Usage

```
Grid.Adjmatrix.Transfer(grid, euclidean.dist=1)
```

Arguments

`grid` a x by y matrix or data.frame, representing y vectors with dimension of x row binded (rbind) together.

`euclidean.dist` a number representing the maximum euclidean distance to be considered as connected. The default setting is 1.

Value

a y by y binary (0/1) adjacency matrix, where y is the row of the argument grid.

Note

Although in real world, the grid coordinates are less than 3-dimension, however, this function is still applicable for vectors larger than 3

Examples

```
###10 3-D coordinates
x <- matrix(rnorm(3*10), nrow = 10)
###a 10x10 adjacency matrix returned for 'net'
###Distance larger than 1 is considered as connected
net<-Grid.Adjmatrix.Transfer(grid=x, euclidean.dist=1)
```

HyperPara.Select *Selecting Hyper Parameters by Bayesian Model Averaging*

Description

This function is aiming at selecting appropriate hyper parameters by Bayesian Model Averaging. This process of this function has already embedded in function `Networks.STD()` and `Networks.Fast()`. The purpose of this function is for having the selected hyper-parameters independently.

Usage

```
HyperPara.Select(net,pvalue,piall,rhoall,n=30)
```

Arguments

<code>net</code>	an "n" by "n" binary adjacent matrix (0/1) for the network configuration with $n = \text{length}(pvalue)$
<code>pvalue</code>	a vector of p-values obtained from large scale statistical hypothesis testing
<code>piall</code>	a vector of possible choices for "pi0" in an increasing order
<code>rhoall</code>	a vector of possible choices of "rho0" and "rho1" in an increasing order
<code>n</code>	a number of iterations you set for Bayesian Model Averaging. The default setting is 30, which is accord with the embedded inner process of in function <code>Networks.STD()</code> and <code>Networks.Fast()</code> . More iterations could be set if you have a desire to have more accurate parameter

Value

a list of selected parameter

pi0 the selected pi0

rho0 the selected rho0

rho1 the selected rho1

Author(s)

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References

Yize Zhao, Jian Kang, Tianwei Yu (2014) A Bayesian nonparametric model for selecting gene and gene sub network, *Annals of Applied Statistics*, in press.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

LikelihoodHistory *Calculating and Plot the history of log-Likelihood value*

Description

This function is a supplementary function for calculating and plotting the history of log-Likelihood value for each iteration. This function is used for tracking the evolution of the MCMC chain obtained from functions `Networks.STD()` and `Networks.Fast()`

Usage

```
LikelihoodHistory(Trace,pvalue,Status=FALSE,True.Node)
```

Arguments

Trace	the trace (results of each iteration) generated by the functions <code>Networks.STD()</code> and <code>Networks.Fast()</code> . Please set as <code>Trace=total\$trace</code> where <code>total</code> is the output of the two functions, <code>Networks.STD()</code> and <code>Networks.Fast()</code>
pvalue	a vector of p-values obtained from large scale statistical hypothesis testing. It should be the same vector which you set for the argument <code>pvalue</code> in functions <code>Networks.STD()</code> and <code>Networks.Fast()</code>
Status	a logical variable indicating whether you have the assumed true value. <code>TRUE</code> means you have the assumed true value for plotting and <code>FALSE</code> , otherwise. The default setting is <code>FALSE</code>
True.Node	a binary(0/1) vector providing the assumed information of each node. The length of vector should be equal to the length of the vector of <code>pvalue</code>

Value

a table indicating the history of the history of log-Likelihood value for each iteration

Note

A plot describing the history of the history of log-Likelihood value for each iteration will be outputted. If you set `Status` and `True.Node`, a red line indicates the log-Likelihood value of the assumed status of each node.

Author(s)

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References

Yize Zhao, Jian Kang, Tianwei Yu (2014) A Bayesian nonparameteric model for selecting gene and gene subnetwork, *Annals of Applied Statistics*, in press.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Networks.Fast

Bayesian Network Discovery using a Hybrid Fast Algorithm

Description

This function implements a hybrid fast algorithm to perform feature selection and sub-network discovery using a Bayesian nonparametric model based on Dirichlet process mixture models, finite mixture of normals model and the Ising model.

Usage

```
Networks.Fast(pvalue, net, iter=5000, nburns=2000,
  algorithms=c("EM", "DPM"),
  v=20, DPM.mcmc=list(nburn=2000, nsave=1, nskip=0, ndisplay=10),
  DPM.prior=list(a0=2, b0=1, m2=rep(0, 1), s2=diag(100000, 1),
  psiinv2=solve(diag(0.5, 1)),
  nu1=4, nu2=4, tau1=1, tau2=100),
  DPparallel=FALSE, n.cores=1, pi.all=c(0.8, 0.85, 0.9, 0.95),
  rho.all=c(1, 2, 5, 10, 15),
  show.steps=10, show.likelihood=FALSE, likelihood.frequency=100)
```

Arguments

pvalue	a vector of p-values obtained from large scale statistical hypothesis testing
net	an "n" by "n" binary (0/1) adjacent matrix of network configurations, where n=length(pvalue)
iter	number of iterations; the default is 5000
nburns	number of burn-in; the default is 2000
algorithms	character taking value "EM" or "DPM" indicating the function to be used to obtain Finite Gaussian Mixture (FGM) estimates. It is recommended to choose "DPM" when the dimension of data is large, and to choose "EM" when the dimension is small.
v	number of iterations set for DPM fitting. v is only valid when you choose algorithms as "DPdensity"
DPM.mcmc	a list giving the MCMC parameters for DPM fitting; see the argument mcmc of function DPdensity() in DPpackage for details; the default setting is DPM.mcmc=list(nburn=2000, nsave=1, nskip=0, ndisplay=10)

DPM.prior	a list giving the prior information; see the argument prior of function DPdensity() in DPPackage for details; the default setting is prior2
piall	a vector of possible choices for "pi0" in an increasing order; the default value is c(0.75, 0.8, 0.85, 0.9)
rhoall	a vector of possible choices of "rho0" and "rho1" in an increasing order; the default value is c(0.5, 1, 5, 10, 15)
DPparallel	the logic variable indicating whether apply parallel computing when you set algorithms="DPM"; the default setting is FALSE
n.cores	number of CUP cores for parallel computing, this argument is only valid when you set algorithms="DPM"; the default setting is 1
show.steps	integer representing the frequency of the results of iterations presented, the default setting is 10. The setting is invalid when trace=FALSE. The setting would not affect the data saved, only for printing
showlikelihood	a logical variable indicating whether to show the log-likelihood value simultaneously. Set TRUE if show the log-likelihood value simultaneously, FALSE, otherwise. FALSE is the default setting
likelihood.frequency	a number representing the frequency showing the log-likelihood value simultaneously. For example, setting likelihood.frequency=100 means showing the log-likelihood value every 100 iterations. The default setting is 100 and it is recommended that do not set a small frequency because it slow down the MCMC chain updating

Details

This function implements a Bayesian nonparametric mixture model for feature selection incorporating network information (Zhao et al., 2014):

- $r_{il} g_i, \boldsymbol{\theta} \sim N(\mu_{g_i}, \sigma_{g_i}),$
- $g_i | z_i=k, \mathbf{q}_k \sim \text{Discrete}(\mathbf{a}_k, \mathbf{q}_k),$
- $\boldsymbol{\theta} \sim G_{0k},$ for g in $\mathbf{a}_k,$
- $\mathbf{q}_k \sim \text{Dirichlet}(\tau_k \mathbf{1}_{\{L_k\}}/L_k),$
- $\boldsymbol{\theta} = \{\boldsymbol{\theta}_g\}_{g \text{ in } \mathbf{a}_0 \text{ and } \mathbf{a}_1}$
- $\boldsymbol{\theta}_g = (\mu_g, \sigma_g)$

where we define

Index $\mathbf{a}_0 = (-L_0+1, -L_0+2, \dots, 0)$, $\mathbf{a}_1 = (1, 2, \dots, L_1)$ and the correspondent probability $\mathbf{q}_0 = (q_{-L_0+1}, q_{-L_0+2}, \dots, q_0)$, $\mathbf{q}_1 = (q_1, q_2, \dots, q_{L_1})$, according to the definition of Discrete($\mathbf{a}_k, \mathbf{b}_k$), for example, $\Pr(g_i=L_0+2) = q_{-L_0+2}$.

Assumption In this algorithm, we assume that "important" features should have larger statics comparing to "unimportant" ones without the loss of generality. In this regard, we set the restriction $\mu_g < \mu_{g+1}$ for $g = -L_0+1, -L_0+2, \dots, L_1$.

This function implements the NET-DPM-3 Zhao et al.(2014). Please refer to the Appendix B.3 for more details.

Value

An object of class "Networks.Fast" containing a list of values, such as, the posterior samples of "z_i" and summary statistics

trace an length(pvalue) by (iter-nburns) matrix

convergence MCMC Heidelberger and Welch convergence diagnostic

graph An igraph graph object of full network

statistics a list of summary statistics characterizing the posterior distribution of "z_i"

mean posterior mean for each feature

median posterior median for each feature

var posterior variance for each feature

quantile posterior quantiles for each feature

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References

Zhao, Y., Kang, J., Yu, T. A Bayesian nonparametric mixture model for selecting gene and gene-sub network, *Annals of Applied Statistics*, In press: 2014.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Examples

```
####Gene Network discovery
##Generating Scale free Gene Network
library(igraph)
g <- barabasi.game(50, power=1, zero.appeal=1.5,directed = FALSE)
net=as(get.adjacency(g,attr=NULL),"matrix")
##Random assign selected genes and make the signal intension as gaussian mixture
newz=rep(c(1,0,0,1,0),10)
Simnorm=function(n){
  weight = c(0.4, 0.6)
  mu = c(8,6)
  sigma = c(1,0.5)
  z = sample(c(1,2),size=n, prob=weight,replace=TRUE)
  r = rnorm(n,mean=mu[z],sd=sigma[z])
  return(r)
}
testcov<-0
for(i in 1:50){
  if(newz[i]!=0){
    testcov[i]<-rnorm(1,mean=0,sd=1)
  }else{
    testcov[i]<-Simnorm(1)
  }
}
```

```

    }
  }
  pvalue=pnorm(-testcov)
  total1=Networks.Fast(pvalue,net,iter=5,nburns=2,
v=20,algorithms="DPM",DPparallel=FALSE,
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(1, 2, 5, 10, 15)
)

```

Networks.STD

Bayesian Network Discovery using a Standard MCMC Algorithm

Description

This function implements the standard Markov chain Monte Carlo (MCMC) algorithm to perform feature selection and sub-network discovery using a Bayesian nonparametric model based on the Dirichlet process mixture (DPM) model and the Ising model.

Usage

```

Networks.STD(pvalue,net,iter=5000, nburns=2000,
piall=c(0.75, 0.8, 0.85, 0.9),rhoall=c(0.5, 1, 5, 10, 15),
status=FALSE,fit,show.steps=1,showlikelihood=FALSE,likelihood.frequency=100)

```

Arguments

pvalue	a vector of p-values obtained from large scale statistical hypothesis testing
net	an "n" by "n" binary adjacent matrix (0/1) for the network configuration with n=length(pvalue)
iter	the number of iterations; the default is 5000
nburns	the number of burn-in; the default is 2000
piall	a vector of possible choices for "pi0" in an increasing order; the default value is c(0.75, 0.8, 0.85, 0.9)
rhoall	a vector of possible choices of "rho0" and "rho1" in an increasing order; the default value is c(0.5, 1, 5, 10, 15)
status	a logical variable indicating whether the you are initial an MCMC chain or resume a chain
fit	a list provide the current values of parameters which are used for resume the chain
show.steps	integer representing the frequency of the results of iterations presented, the default value is 1.The setting is invalid when trace=FALSE. The setting would not affect the data saved, only for printing
showlikelihood	a logical variable indicating whether to show the log-likelihood value simultaneously.Set TRUE if show the log-likelihood value simultaneously, FALSE, otherwise. FALSE is the default setting

likelihood.frequency

a number representing the frequency showing the log-likelihood value simultaneously. For example, setting likelihood.frequency=100 means showing the log-likelihood value every 100 iterations. The default setting is 100 and it is recommended that do not set a small frequency because it slow down the MCMC chain updating

Details

This function implements a Bayesian nonparametric mixture model for feature selection incorporating network information (Zhao et al., 2014):

- $r_{ij} | g_i, \boldsymbol{\theta}$ $\sim N(\mu_{g_i}, \sigma_{g_i})$,
- $g_i | z_i=k, \mathbf{q}_k \sim \text{Discrete}(\mathbf{a}_k, \mathbf{q}_k)$,
- $\boldsymbol{\theta} \sim G_{0k}$, for g in \mathbf{a}_k ,
- $\mathbf{q}_k \sim \text{Dirichlet}(\tau_k \mathbf{1}_{\{L_k\}}/L_k)$,
- $\boldsymbol{\theta} = \{\boldsymbol{\theta}_g\}_{g \text{ in } \mathbf{a}_0 \text{ and } \mathbf{a}_1}$
- $\boldsymbol{\theta}_g = (\mu_g, \sigma_g)$

where we define

Index $\mathbf{a}_0 = (-L_0+1, -L_0+2, \dots, 0)$, $\mathbf{a}_1 = (1, 2, \dots, L_1)$ and the correspondent probability $\mathbf{q}_0 = (q_{-L_0+1}, q_{-L_0+2}, \dots, q_0)$, $\mathbf{q}_1 = (q_1, q_2, \dots, q_{L_1})$, according to the definition of $\text{Discrete}(\mathbf{a}_k, \mathbf{b}_k)$, for example, $\Pr(g_i=L_0+2) = q_{-L_0+2}$.

Assumption In this algorithm, we assume that "important" features should have larger statistics comparing to "unimportant" ones without the loss of generality. In this regard, we set the restriction $\mu_g < \mu_{g+1}$ for $g = -L_0+1, -L_0+2, \dots, L_1$.

This function implements the NET-DPM-1 Zhao et al.(2014). Please refer to the Appendix B.1 for more details.

Value

An object of class "Networks.STD" containing Bayesian information of the MCMC chain

trace a matrix of dimension (iter-nburns) by length(pvalue) containing posterior samples of classification label "g_i"

convergence MCMC Heidelberger and Welch convergence diagnostic

graph An igraph graph object of full network

model a list containing parameters of the MCMC chain for resuming the chain

Author(s)

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References

Zhao, Y., Kang, J., Yu, T. A Bayesian nonparametric mixture model for selecting gene and gene-sub network, *Annals of Applied Statistics*, In press: 2014.

Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Examples

```
library(igraph)
###Creating the network of 10X10 image
g <- graph.lattice(length=10,dim=2)
net=as(get.adjacency(g,attr=NULL),"matrix")##this is the input of argument 'net'
##Assign the signal elements with signal intention
##as normal distribution N(1,0.2). While noise is set as N(0,0.2)
newz=rep(0,100)
for (i in 3:7)
{
  newz[(i*10+3):(i*10+7)]=1
}
testcov<-0
for(i in 1:100){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=0.2)

  }else{
    testcov[i]<-rnorm(1,mean=1,sd=0.2)

  }
}
##The profile of the image
image(matrix(testcov,10,10),col=gray(seq(0,1,length=255)))
##Transform the signals into pvalue form and begin identification
pvalue=pnorm(-testcov)
total=Networks.STD(pvalue,net,iter=3,nburns=1,
pia11=c(0.8, 0.85, 0.9, 0.95),rhoall=c(0.5,1,5,10,15))
```

plot.Networks.Fast *Plotting Bayesian Network Discovery using a Hybrid Fast Algorithm*

Description

plot method for class "Networks.Fast"

Usage

```
## S3 method for class 'Networks.Fast'
plot(x, ...)
```

Arguments

x An object of class "Networks.Fast" resulting of a call to [Networks.Fast](#).
 ... other arguments

See Also

[Networks.Fast](#).

Examples

```
library(igraph)
####Gene Network discovery
##Generating Scale free Gene Network
library(igraph)
g <- barabasi.game(50, power=1, zero.appeal=1.5,directed = FALSE)
net=as(get.adjacency(g,attr=NULL),"matrix")
##Random assign selected genes and make the signal intension as gaussian mixture
newz=rep(c(1,0,0,1,0),10)
Simnorm=function(n){
  weight = c(0.4, 0.6)
  mu = c(8,6)
  sigma = c(1,0.5)
  z = sample(c(1,2),size=n, prob=weight,replace=TRUE)
  r = rnorm(n,mean=mu[z],sd=sigma[z])
  return(r)
}
testcov<-0
for(i in 1:50){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=1)
  }else{
    testcov[i]<-Simnorm(1)
  }
}
pvalue=pnorm(-testcov)
total1=Networks.Fast(pvalue,net,iter=5,nburns=2,
v=20,algorithms="DPM",DPparallel=FALSE,
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(1, 2, 5, 10, 15)
)

#####Plotting the object of class "Networks.Fast"
plot(total1)
```

plot.Networks.STD

Plotting Bayesian Network Discovery using a Standard MCMC Algorithm

Description

plot method for class "Networks.STD"

Usage

```
## S3 method for class 'Networks.STD'
plot(x, ...)
```

Arguments

x An object of class "Networks.STD" resulting of a call to [Networks.STD](#).

... other arguments

See Also

[Networks.STD](#).

Examples

```
library(igraph)
###Creating the network of 10X10 image
g <- graph.lattice(length=10,dim=2)
net=as(get.adjacency(g,attr=NULL),"matrix")##this is the input of argument 'net'
##Assign the signal elements with signal intention
##as normal distribution N(1,0.2). While noise is set as N(0,0.2)
newz=rep(0,100)
for (i in 3:7)
{
  newz[(i*10+3):(i*10+7)]=1
}
testcov<-0
for(i in 1:100){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=0.2)

  }else{
    testcov[i]<-rnorm(1,mean=1,sd=0.2)

  }
}
##The profile of the image
image(matrix(testcov,10,10),col=gray(seq(0,1,length=255)))
##Transform the signals into pvalue form and begin identification
pvalue=pnorm(-testcov)
total=Networks.STD(pvalue,net,iter=3,nburns=1,
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(0.5,1,5,10,15))

#####plot the object of class "Networks.STD"
plot(total)
```

Plot.Subnetwork	<i>Plotting the Network Feature Selected</i>
-----------------	--

Description

This is a supportive function for plotting the network feature according to the outputs of `Network.STD` and `Network.Fast`. If the mean of the marginal distribution of a certain node is larger than 0.5, we consider this node as selected.

Usage

```
Plot.Subnetwork(net, trace)
```

Arguments

net	an "n" by "n" binary (0/1) adjacent matrix of network configurations, where $n = \text{length}(pvalue)$; same as the argument net in function <code>Network.STD()</code> and <code>Network.Fast()</code>
trace	the trace (results of each iteration) generated by the functions <code>Networks.STD()</code> and <code>Networks.Fast()</code> . Please set as <code>trace=total\$trace</code> where total is the output of the two functions, <code>Networks.STD()</code> and <code>Networks.Fast()</code>

Value

A plot of the network feature selected

Examples

```
library(igraph)
g=barabasi.game(10,power=0.5)
net=as(get.adjacency(g),"matrix")
trace=matrix(c(rep(1,5),rep(0,5)),nrow=1)
Plot.Subnetwork(net,trace)
```

Subnetwork.Select	<i>Summarize the information of the sub networks selected by Network.Fast() and Network.STD()</i>
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Description

This function is to Summarize the information of the sub networks selected by `Network.Fast()` and `Network.STD()`. If the marginal distribution of a certain node is larger than 0.5, we consider this node as selected. The aim of this function is to customize the process of summarizing.

Usage

```
Subnetwork.Select(net, trace, node.based=NULL, infinite=TRUE, steps=5)
```

Arguments

net	an "n" by "n" binary (0/1) adjacent matrix of network configurations, where $n = \text{length}(pvalue)$; same as the argument <code>net</code> in function <code>Network.STD</code> and <code>Network.Fast</code>
trace	the trace (results of each iteration) generated by the functions <code>Networks.STD()</code> and <code>Networks.Fast()</code> . Please set as <code>trace=total\$trace</code> where <code>total</code> is the output of the two functions, <code>Networks.STD()</code> and <code>Networks.Fast()</code> ; you can also provide a binary 1 by "n" matrix representing the "important"(1) and "unimportant"(0) nodes
node.based	a vector representing the orders of the nodes the sub network centered
infinite	a logical variable indicating whether the expanding of sub network in a "infinite" pattern or not; <code>infinite=TRUE</code> if searching the "important" sub network based on one node without the limitation of maximum steps; <code>infinite=FALSE</code> if searching the sub network under a regulated way
steps	a number representing the maximum steps for expanding the sub network

Value

a list of values containing the the information of the sub networks selected by `Network.Fast()` and `Network.STD()`

eids a vector of order representing the selected nodes

adj a "m" by "m" binary adjacent matrix of the sub network configurations where $n = \text{length}(eids)$

summary.Networks.Fast *Summarizing Bayesian Network Discovery using a Hybrid Fast Algorithm*

Description

Summary method for class "Networks.Fast"

Usage

```
## S3 method for class 'Networks.Fast'
summary(object, ...)
```

Arguments

object	An object of class "Networks.Fast" resulting of a call to Networks.Fast .
...	other arguments

See Also

[Networks.Fast.](#)

Examples

```

library(igraph)
####Gene Network discovery
##Generating Scale free Gene Network
library(igraph)
g <- barabasi.game(50, power=1, zero.appeal=1.5,directed = FALSE)
net=as(get.adjacency(g,attr=NULL),"matrix")
##Random assign selected genes and make the signal intension as gaussian mixture
newz=rep(c(1,0,0,1,0),10)
Simnorm=function(n){
weight = c(0.4, 0.6)
mu = c(8,6)
sigma = c(1,0.5)
z = sample(c(1,2),size=n, prob=weight,replace=TRUE)
r = rnorm(n,mean=mu[z],sd=sigma[z])
return(r)
}
testcov<-0
for(i in 1:50){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=1)
  }else{
    testcov[i]<-Simnorm(1)
  }
}
pvalue=pnorm(-testcov)
total1=Networks.Fast(pvalue,net,iter=5,nburns=2,
v=20,algorithms="DPM",DPparallel=FALSE,
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(1, 2, 5, 10, 15)
)

#####summary the object of class "Networks.Fast"
summary(total1)

```

summary.Networks.STD *Summarizing Bayesian Network Discovery using a Standard MCMC Algorithm*

Description

Summary method for class "Networks.STD"

Usage

```

## S3 method for class 'Networks.STD'
summary(object, ...)

```

Arguments

object An object of class "Networks.STD" resulting of a call to [Networks.STD](#).
 ... other arguments

See Also

[Networks.STD](#).

Examples

```
library(igraph)
###Creating the network of 10X10 image
g <- graph.lattice(length=10,dim=2)
net=as(get.adjacency(g,attr=NULL),"matrix")##this is the input of argument 'net'
##Assign the signal elements with signal intention
##as normal distribution N(1,0.2). While noise is set as N(0,0.2)
newz=rep(0,100)
for (i in 3:7)
{
  newz[(i*10+3):(i*10+7)]=1
}
testcov<-0
for(i in 1:100){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=0.2)

  }else{
    testcov[i]<-rnorm(1,mean=1,sd=0.2)

  }
}
##The profile of the image
image(matrix(testcov,10,10),col=gray(seq(0,1,length=255)))
##Transform the signals into pvalue form and begin identification
pvalue=pnorm(-testcov)
total=Networks.STD(pvalue,net,iter=3,nburns=1,
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(0.5,1,5,10,15))

#####summary the object of class "Networks.STD"
summary(total)
```

Description

This function is to summarize the accuracy of node and Network selection achieved by the functions `Networks.STD` and `Networks.Fast`. This function is designed to validate whether the settings of arguments are appropriate for one simulation study and provide insights of setting arguments for real data.

Usage

```
SummaryAccuracy(Trace, No.Sets,
Type.Accuracy=c("Node", "Sub-network"), True.Node,
TruePositive.Net, FalsePositive.Net,
Type.Net.Accuracy=c("Marginal", "Sample"), Tolerance)
```

Arguments

Trace	the trace (results of each iteration) generated by the functions <code>Networks.STD()</code> and <code>Networks.Fast()</code> . For <code>Networks.STD()</code> and <code>Networks.Fast()</code> , set <code>Trace=total\$trace</code> where <code>total</code> is the output of the two functions, <code>Networks.STD()</code> and <code>Networks.Fast()</code> . Use <code>rbind()</code> bind the inputs if you have multiple sets of results
No.Sets	number of sets of results you put in the argument Trace
Type.Accuracy	character taking value "Node" or "Sub-network". "Node" represents checking the accuracy of each node; "Sub-network" represents checking the accuracy of network of interest
True.Node	a binary(0/1) vector providing the information of each node. The length of vector should be equal to the length of the vector of <code>pvalue</code>
TruePositive.Net	a vector representing the network of interest consisted of assumed positive nodes. For example, if you assume No.6, No.8 and No.9 nodes consist the network of interest, set <code>TruePositive.Net=c(6, 8, 9)</code>
FalsePositive.Net	a vector representing the assumed negative nodes which would lead false positive networking (larger network) if such nodes are assumed positive in the results. Such nodes should be connected to the network of interest and assumed negative. For example, if you assume No.3 and No.11 nodes would lead false positive networking (larger network), set <code>TruePositive.Net=c(3, 11)</code>
Type.Net.Accuracy	character taking value "Marginal" or "Sample". "Marginal" represents using marginal distribution of each data-set for checking accuracy, for example, if the probability (mean) of one node of a data-set is larger than 0.5, we consider it positive, otherwise, negative. Setting "Sample" represents checking accuracy through each iteration. For <code>Type.Accuracy="Node"</code> , only when <code>Type.Net.Accuracy="Sample"</code> , is valid.
Tolerance	percentage of the Tolerance rate of accuracy. This argument is only valid when set <code>Type.Accuracy="Networking"</code> . Setting <code>Tolerance=0.9</code> means we consider selecting a correct networking selection if more than 0.9 positive nodes of networking or <code>TruePositive.Net</code> are considered as positive AND less than 0.1 negative nodes of networking or <code>FalsePositive.Net</code> are considered as positive.

Value

a list of values containing the information of accuracy of nodes and network selection

TPR.average The Average of True Positive Rate

FPR.average The Average of True False Rate
FDR.average The Average of False Positive Rate
TPR True Positive Rate of each node
FPR False Positive Rate of each node

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References

Yize Zhao, Jian Kang, Tianwei Yu (2014) A Bayesian nonparametric model for selecting gene and gene sub network, Annals of Applied Statistics, in press.
 Zhou Lan, Jian Kang, Tianwei Yu, Yize Zhao, BANFF: an R package for network identifications via Bayesian nonparametric mixture models, working paper.

Examples

```
####Gene Network discovery
##Generating Scale free Gene Network
library(igraph)
library(BANFF)
g <- barabasi.game(50, power=1, zero.appeal=1.5, directed = FALSE)
net=as(get.adjacency(g,attr=NULL),"matrix")
##Random assign selected genes and make the signal intension as gaussian mixture
newz=rep(c(1,0,0,1,0),10)
Simnorm=function(n){
  weight = c(0.4, 0.6)
  mu = c(8,6)
  sigma = c(1,0.5)
  z = sample(c(1,2),size=n, prob=weight,replace=TRUE)
  r = rnorm(n,mean=mu[z],sd=sigma[z])
  return(r)
}
testcov<-0
for(i in 1:50){
  if(newz[i]==0){
    testcov[i]<-rnorm(1,mean=0,sd=1)
  }else{
    testcov[i]<-Simnorm(1)
  }
}
pvalue=pnorm(-testcov)
total1=Networks.Fast(pvalue,net,iter=5,nburns=2,
v=20,algorithms="DPM",
piall=c(0.8, 0.85, 0.9, 0.95),rhoall=c(1, 2, 5, 10, 15))
####checking accuracy
##checking accuracy of nodes
SummaryAccuracy(Trace=total1$trace,No.Sets=1,Type.Accuracy="Node",
True.Node=newz,Type.Net.Accuracy="Sample")
```

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