

Package ‘TensorClustering’

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Type Package

Title Model-Based Tensor Clustering

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Description Performs model-based tensor clustering methods including Tensor Gaussian Mixture Model (TGMM), Tensor Envelope Mixture Model (TEMM) by Deng and Zhang (2021) <[DOI:10.1111/biom.13486](https://doi.org/10.1111/biom.13486)>, Doubly-Enhanced EM (DEEM) algorithm by Mai, Zhang, Pan and Deng (2021) <[DOI:10.1080/01621459.2021.1904959](https://doi.org/10.1080/01621459.2021.1904959)>.

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Contents

DEEM	2
TEMM	4
TGMM	6
tune_K	8
tune_lamb	9
tune_u_joint	11
tune_u_sep	12

DEEM	<i>Doubly-enhanced EM algorithm</i>
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Description

Doubly-enhanced EM algorithm for tensor clustering

Usage

```
DEEM(X, nclass, niter = 100, lambda = NULL, dfmax = n, pmax = nvars, pf = rep(1, nvars),
     eps = 1e-04, maxit = 1e+05, sm1 = 1e-06, verbose = FALSE, ceps = 0.1,
     initial = TRUE, vec_x = NULL)
```

Arguments

<code>X</code>	Input tensor (or matrix) list of length n , where n is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any positive integer not less than 2.
<code>nclass</code>	Number of clusters.
<code>niter</code>	Maximum iteration times for EM algorithm. Default value is 100.
<code>lambda</code>	A user-specified lambda value. lambda is the weight of L1 penalty and a smaller lambda allows more variables to be nonzero
<code>dfmax</code>	The maximum number of selected variables in the model. Default is the number of observations n .
<code>pmax</code>	The maximum number of potential selected variables during iteration. In middle step, the algorithm can select at most $pmax$ variables and then shrink part of them such that the number of final selected variables is less than $dfmax$.
<code>pf</code>	Weight of lasso penalty. Default is a vector of value 1 and length p , representing L1 penalty of length p . Can be modified to use adaptive lasso penalty.
<code>eps</code>	Convergence threshold for coordinate descent difference between iterations. Default value is $1e-04$.
<code>maxit</code>	Maximum iteration times for coordinate descent for all lambda. Default value is $1e+05$.
<code>sm1</code>	Threshold for ratio of loss function change after each iteration to old loss function value. Default value is $1e-06$.
<code>verbose</code>	Indicates whether print out lambda during iteration or not. Default value is FALSE.
<code>ceps</code>	Convergence threshold for cluster mean difference between iterations. Default value is 1.
<code>initial</code>	Whether to initialize algorithm with K-means clustering. Default value is TRUE.
<code>vec_x</code>	Vectorized tensor data. Default value is NULL

Details

The `DEEM` function implements the Doubly-Enhanced EM algorithm (DEEM) for tensor clustering. The observations \mathbf{X}_i are assumed to be following the tensor normal mixture model (TNMM) with common covariances across different clusters:

$$\mathbf{X}_i \sim \sum_{k=1}^K \pi_k \text{TN}(\boldsymbol{\mu}_k; \boldsymbol{\Sigma}_1, \dots, \boldsymbol{\Sigma}_M), \quad i = 1, \dots, n,$$

where $0 < \pi_k < 1$ is the prior probability for \mathbf{X} to be in the k -th cluster such that $\sum_{k=1}^K \pi_k = 1$, $\boldsymbol{\mu}_k$ is the cluster mean of the k -th cluster and $\boldsymbol{\Sigma}_1, \dots, \boldsymbol{\Sigma}_M$ are the common covariances across different clusters. Under the TNMM framework, the optimal clustering rule can be showed as

$$\hat{Y}^{opt} = \arg \max_k \{ \log \pi_k + \langle \mathbf{X} - (\boldsymbol{\mu}_1 + \boldsymbol{\mu}_k)/2, \mathbf{B}_k \rangle \},$$

where $\mathbf{B}_k = \llbracket \boldsymbol{\mu}_k - \boldsymbol{\mu}_1; \boldsymbol{\Sigma}_1^{-1}, \dots, \boldsymbol{\Sigma}_M^{-1} \rrbracket$. In the enhanced E-step, `DEEM` imposes sparsity directly on the optimal clustering rule as a flexible alternative to popular low-rank assumptions on tensor coefficients \mathbf{B}_k as

$$\min_{\mathbf{B}_2, \dots, \mathbf{B}_K} \left[\sum_{k=2}^K (\langle \mathbf{B}_k, \llbracket \hat{\boldsymbol{\Sigma}}_1^{(t)}, \dots, \hat{\boldsymbol{\Sigma}}_M^{(t)} \rrbracket \rangle - 2 \langle \mathbf{B}_k, \hat{\boldsymbol{\mu}}_k^{(t)} - \hat{\boldsymbol{\mu}}_1^{(t)} \rangle) + \lambda^{(t+1)} \sum_{\mathcal{J}} \sqrt{\sum_{k=2}^K b_{k,\mathcal{J}}^2} \right],$$

where $\lambda^{(t+1)}$ is a tuning parameter. In the enhanced M-step, `DEEM` employs a new estimator for the tensor correlation structure, which facilitates both the computation and the theoretical studies.

Value

<code>pi</code>	A vector of estimated prior probabilities for clusters.
<code>mu</code>	A list of estimated cluster means.
<code>sigma</code>	A list of estimated covariance matrices.
<code>gamma</code>	A n by <code>nclass</code> matrix of estimated membership weights.
<code>y</code>	A vector of estimated labels.
<code>iter</code>	Number of iterations until convergence.
<code>df</code>	Average zero elements in beta over iterations.
<code>beta</code>	A matrix of vectorized \mathbf{B}_k .

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

Mai, Q., Zhang, X., Pan, Y. and Deng, K. (2021). A Doubly-Enhanced EM Algorithm for Model-Based Tensor Clustering. *Journal of the American Statistical Association*.

See Also

[tune_lamb](#), [tune_K](#)

Examples

```

dimen = c(5,5,5)
nvars = prod(dimen)
K = 2
n = 100
sigma = array(list(),3)

sigma[[1]] = sigma[[2]] = sigma[[3]] = diag(5)

B2=array(0,dim=dimen)
B2[1:3,1,1]=2

y = c(rep(1,50),rep(2,50))
M = array(list(),K)
M[[1]] = array(0,dim=dimen)
M[[2]] = B2

vec_x=matrix(rnorm(n*prod(dimen)),ncol=n)
X=array(list(),n)
for (i in 1:n){
  X[[i]] = array(vec_x[,i],dim=dimen)
  X[[i]] = M[[y[i]]] + X[[i]]
}

myfit = DEEM(X, nclass=2, lambda=0.05)

```

 TEMM

Fit the Tensor Envelope Mixture Model (TEMM)

Description

Fit the Tensor Envelope Mixture Model (TEMM)

Usage

```

TEMM(Xn, u, K, initial = "kmeans", iter.max = 500,
stop = 1e-3, trueY = NULL, print = FALSE)

```

Arguments

Xn	The tensor for clustering, should be array type, the last dimension is the sample size n.
u	A vector of envelope dimension
K	Number of clusters, greater than or equal to 2.
initial	Initialization method for the regularized EM algorithm. Default value is "kmeans".
iter.max	Maximum number of iterations. Default value is 500.

stop	Convergence threshold of relative change in cluster means. Default value is 1e-3.
trueY	A vector of true cluster labels of each observation. Default value is NULL.
print	Whether to print information including current iteration number, relative change in cluster means and clustering error (%) in each iteration.

Details

The TEMM function fits the Tensor Envelope Mixture Model (TEMM) through a subspace-regularized EM algorithm. For mode m , let $(\mathbf{\Gamma}_m, \mathbf{\Gamma}_{0m}) \in R^{p_m \times p_m}$ be an orthogonal matrix where $\mathbf{\Gamma}_m \in R^{p_m \times u_m}$, $u_m \leq p_m$, represents the material part. Specifically, the material part $\mathbf{X}_{*,m} = \mathbf{X} \times_m \mathbf{\Gamma}_m^T$ follows a tensor normal mixture distribution, while the immaterial part $\mathbf{X}_{\circ,m} = \mathbf{X} \times_m \mathbf{\Gamma}_{0m}^T$ is unimodal, independent of the material part and hence can be eliminated without loss of clustering information. Dimension reduction is achieved by focusing on the material part $\mathbf{X}_{*,m} = \mathbf{X} \times_m \mathbf{\Gamma}_m^T$. Collectively, the joint reduction from each mode is

$$\mathbf{X}_* = \llbracket \mathbf{X}; \mathbf{\Gamma}_1^T, \dots, \mathbf{\Gamma}_M^T \rrbracket \sim \sum_{k=1}^K \pi_k \text{TN}(\boldsymbol{\alpha}_k; \boldsymbol{\Omega}_1, \dots, \boldsymbol{\Omega}_M), \quad \mathbf{X}_* \perp\!\!\!\perp \mathbf{X}_{\circ,m},$$

where $\boldsymbol{\alpha}_k \in R^{u_1 \times \dots \times u_M}$ and $\boldsymbol{\Omega}_m \in R^{u_m \times u_m}$ are the dimension-reduced clustering parameters and $\mathbf{X}_{\circ,m}$ does not vary with cluster index Y . In the E-step, the membership weights are evaluated as

$$\hat{\eta}_{ik}^{(s)} = \frac{\hat{\pi}_k^{(s-1)} f_k(\mathbf{X}_i; \hat{\boldsymbol{\theta}}^{(s-1)})}{\sum_{k=1}^K \hat{\pi}_k^{(s-1)} f_k(\mathbf{X}_i; \hat{\boldsymbol{\theta}}^{(s-1)})},$$

where f_k denotes the conditional probability density function of \mathbf{X}_i within the k -th cluster. In the subspace-regularized M-step, the envelope subspace is iteratively estimated through a Grassmann manifold optimization that minimize the following log-likelihood-based objective function:

$$G_m^{(s)}(\mathbf{\Gamma}_m) = \log |\mathbf{\Gamma}_m^T \mathbf{M}_m^{(s)} \mathbf{\Gamma}_m| + \log |\mathbf{\Gamma}_m^T (\mathbf{N}_m^{(s)})^{-1} \mathbf{\Gamma}_m|,$$

where $\mathbf{M}_m^{(s)}$ and $\mathbf{N}_m^{(s)}$ are given by

$$\mathbf{M}_m^{(s)} = \frac{1}{np-m} \sum_{i=1}^n \sum_{k=1}^K \hat{\eta}_{ik}^{(s)} (\boldsymbol{\epsilon}_{ik}^{(s)})_{(m)} (\hat{\boldsymbol{\Sigma}}_{-m}^{(s-1)})^{-1} (\boldsymbol{\epsilon}_{ik}^{(s)})_{(m)}^T,$$

$$\mathbf{N}_m^{(s)} = \frac{1}{np-m} \sum_{i=1}^n (\mathbf{X}_i)_{(m)} (\hat{\boldsymbol{\Sigma}}_{-m}^{(s-1)})^{-1} (\mathbf{X}_i)_{(m)}^T.$$

The intermediate estimators $\mathbf{M}_m^{(s)}$ can be viewed the mode- m conditional variation estimate of $\mathbf{X} | Y$ and $\mathbf{N}_m^{(s)}$ is the mode- m marginal variation estimate of \mathbf{X} .

Value

id	A vector of estimated labels.
pi	A vector of estimated prior probabilities for clusters.
eta	A n by K matrix of estimated membership weights.

<code>Mu.est</code>	A list of estimated cluster means.
<code>SIG.est</code>	A list of estimated covariance matrices.
<code>Mm</code>	Estimation of M_m defined in paper.
<code>Nm</code>	Estimation of N_m defined in paper.
<code>Gamma.est</code>	A list of estimated envelope basis.
<code>PGamma.est</code>	A list of envelope projection matrices.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

Deng, K. and Zhang, X. (2021). Tensor Envelope Mixture Model for Simultaneous Clustering and Multiway Dimension Reduction. *Biometrics*.

See Also

[TGMM](#), [tune_u_sep](#), [tune_u_joint](#)

Examples

```
A = array(c(rep(1,20),rep(2,20))+rnorm(40),dim=c(2,2,10))
myfit = TEMM(A,u=c(2,2),K=2)
```

TGMM

Fit the Tensor Gaussian Mixture Model (TGMM)

Description

Fit the Tensor Gaussian Mixture Model (TGMM)

Usage

```
TGMM(Xn, K, shape = "shared", initial = "kmeans",
iter.max = 500, stop = 1e-3, trueY = NULL, print = FALSE)
```

Arguments

<code>Xn</code>	The tensor for clustering, should be array type, the last dimension is the sample size n .
<code>K</code>	Number of clusters, greater than or equal to 2.
<code>shape</code>	"shared" if assume common covariance across mixtures, "distinct" if allow different covariance structures. Default value is "shared".
<code>initial</code>	Initialization method for the regularized EM algorithm. Default value is "kmeans".

<code>iter.max</code>	Maximum number of iterations. Default value is 500.
<code>stop</code>	Convergence threshold of relative change in cluster means. Default value is 1e-3.
<code>trueY</code>	A vector of true cluster labels of each observation. Default value is NULL.
<code>print</code>	Whether to print information including current iteration number, relative change in cluster means and clustering error (%) in each iteration.

Details

The `TGMM` function fits the Tensor Gaussian Mixture Model (TGMM) through the classical EM algorithm. TGMM assumes the following tensor normal mixture distribution of M-way tensor data \mathbf{X} :

$$\mathbf{X} \sim \sum_{k=1}^K \pi_k \text{TN}(\boldsymbol{\mu}_k, \mathcal{M}_k), \quad i = 1, \dots, n,$$

where $0 < \pi_k < 1$ is the prior probability for \mathbf{X} to be in the k -th cluster such that $\sum_{k=1}^K \pi_k = 1$, $\boldsymbol{\mu}_k$ is the mean of the k -th cluster, $\mathcal{M}_k \equiv \{\boldsymbol{\Sigma}_{km}, m = 1, \dots, M\}$ is the set of covariances of the k -th cluster. If \mathcal{M}_k 's are the same for $k = 1, \dots, K$, call `TGMM` with argument `shape="shared"`.

Value

<code>id</code>	A vector of estimated labels.
<code>pi</code>	A vector of estimated prior probabilities for clusters.
<code>eta</code>	A n by K matrix of estimated membership weights.
<code>Mu.est</code>	A list of estimated cluster means.
<code>SIG.est</code>	A list of estimated covariance matrices.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

- Deng, K. and Zhang, X. (2021). Tensor Envelope Mixture Model for Simultaneous Clustering and Multiway Dimension Reduction. *Biometrics*.
- Tait, P. A. and McNicholas, P. D. (2019). Clustering higher order data: Finite mixtures of multidimensional arrays. *arXiv:1907.08566*.

See Also

[TEMM](#)

Examples

```
A = array(c(rep(1,20),rep(2,20))+rnorm(40),dim=c(2,2,10))
myfit = TGMM(A,K=2,shape="shared")
```

tune_K	<i>Select the number of clusters K in DEEM</i>
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Description

Select the number of clusters K along with tuning parameter lambda through BIC in DEEM.

Usage

```
tune_K(X, seqK, seqlamb, initial = TRUE, vec_x = NULL)
```

Arguments

X	Input tensor (or matrix) list of length n , where n is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any positive integer not less than 2.
seqK	A sequence of user-specified number of clusters.
seqlamb	A sequence of user-specified lambda values. lambda is the weight of L1 penalty and a smaller lambda allows more variables to be nonzero
initial	Whether to initialize algorithm with K-means clustering. Default value is TRUE.
vec_x	Vectorized tensor data. Default value is NULL

Details

The `tune_K` function runs `tune_lamb` function $\text{length}(\text{seqK})$ times to choose the tuning parameter λ and number of clusters K simultaneously. Let $\hat{\boldsymbol{\theta}}^{\{\lambda, K\}}$ be the output of `DEEM` with the tuning parameter and number of clusters fixed at λ and K respectively, `tune_K` looks for the values of λ and K that minimizes

$$\text{BIC}(\lambda, K) = -2 \sum_{i=1}^n \log\left(\sum_{k=1}^K \hat{\pi}_k^{\{\lambda, K\}} f_k(\mathbf{X}_i; \hat{\boldsymbol{\theta}}_k^{\{\lambda, K\}})\right) + \log(n) \cdot |\hat{\mathcal{D}}^{\{\lambda, K\}}|,$$

where $\hat{\mathcal{D}}^{\{\lambda, K\}} = \{(k, \mathcal{J}) : \hat{b}_{k, \mathcal{J}}^\lambda \neq 0\}$ is the set of nonzero elements in $\hat{\mathbf{B}}_2^{\{\lambda, K\}}, \dots, \hat{\mathbf{B}}_K^{\{\lambda, K\}}$. The `tune_K` function intrinsically selects the initial point and return the optimal estimated labels.

Value

opt_K	Selected number of clusters that leads to optimal BIC.
opt_lamb	Tuned lambda that leads to optimal BIC.
Krank	A selection summary.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

Mai, Q., Zhang, X., Pan, Y. and Deng, K. (2021). A Doubly-Enhanced EM Algorithm for Model-Based Tensor Clustering. *Journal of the American Statistical Association*.

See Also

[DEEM](#), [tune_lamb](#)

Examples

```

dimen = c(5,5,5)
nvars = prod(dimen)
K = 2
n = 100
sigma = array(list(),3)

sigma[[1]] = sigma[[2]] = sigma[[3]] = diag(5)

B2=array(0,dim=dimen)
B2[1:3,1,1]=2

y = c(rep(1,50),rep(2,50))
M = array(list(),K)
M[[1]] = array(0,dim=dimen)
M[[2]] = B2

vec_x=matrix(rnorm(n*prod(dimen)),ncol=n)
X=array(list(),n)
for (i in 1:n){
  X[[i]] = array(vec_x[,i],dim=dimen)
  X[[i]] = M[[y[i]]] + X[[i]]
}

mytune = tune_K(X, seqK=2:4, seqlamb=seq(0.01,0.1,by=0.01))

```

tune_lamb

Parameter tuning in enhanced E-step in DEEM

Description

Perform parameter tuning through BIC in DEEM.

Usage

```
tune_lamb(X, K, seqlamb, initial = TRUE, vec_x = NULL)
```

Arguments

X	Input tensor (or matrix) list of length n , where n is the number of observations. Each element of the list is a tensor or matrix. The order of tensor can be any positive integer not less than 2.
K	Number of clusters.
seq_lamb	A sequence of user-specified lambda values. lambda is the weight of L1 penalty and a smaller lambda allows more variables to be nonzero
initial	Whether to initialize algorithm with K-means clustering. Default value is TRUE.
vec_x	Vectorized tensor data. Default value is NULL

Details

The `tune_lamb` function adopts a BIC-type criterion to select the tuning parameter λ in the enhanced E-step. Let $\hat{\theta}^\lambda$ be the output of `DEEM` with the tuning parameter fixed at λ , `tune_lamb` looks for the value of λ that minimizes

$$\text{BIC}(\lambda) = -2 \sum_{i=1}^n \log\left(\sum_{k=1}^K \hat{\pi}_k^\lambda f_k(\mathbf{X}_i; \hat{\theta}_k^\lambda)\right) + \log(n) \cdot |\hat{\mathcal{D}}^\lambda|,$$

where $\hat{\mathcal{D}}^\lambda = \{(k, \mathcal{J}) : \hat{b}_{k, \mathcal{J}}^\lambda \neq 0\}$ is the set of nonzero elements in $\hat{B}_2^\lambda, \dots, \hat{B}_K^\lambda$. The `tune_lamb` function intrinsically selects the initial point and return the optimal estimated labels.

Value

opt_lamb	Tuned lambda that leads to optimal BIC.
opt_bic	BIC value.
opt_y	Estimated labels fitted by DEEM with tuned lambda.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

Mai, Q., Zhang, X., Pan, Y. and Deng, K. (2021). A Doubly-Enhanced EM Algorithm for Model-Based Tensor Clustering. *Journal of the American Statistical Association*.

See Also

[DEEM](#), [tune_K](#)

Examples

```
dimen = c(5,5,5)
nvars = prod(dimen)
K = 2
n = 100
sigma = array(list(),3)
```

```

sigma[[1]] = sigma[[2]] = sigma[[3]] = diag(5)

B2=array(0,dim=dimen)
B2[1:3,1,1]=2

y = c(rep(1,50),rep(2,50))
M = array(list(),K)
M[[1]] = array(0,dim=dimen)
M[[2]] = B2

vec_x=matrix(rnorm(n*prod(dimen)),ncol=n)
X=array(list(),n)
for (i in 1:n){
  X[[i]] = array(vec_x[,i],dim=dimen)
  X[[i]] = M[[y[i]]] + X[[i]]
}

mytune = tune_lamb(X, K=2, seqlamb=seq(0.01,0.1,by=0.01))

```

tune_u_joint

Tuning envelope dimension jointly by BIC in TEMM.

Description

Tuning envelope dimension jointly by BIC in TEMM.

Usage

```
tune_u_joint(u_candi, K, X, iter.max = 500, stop = 0.001, trueY = NULL)
```

Arguments

u_candi	A list of length M containing candidate envelope dimension for each mode.
K	Number of clusters, greater than or equal to 2.
X	The tensor for clustering, should be array type, the last dimension is the sample size n.
iter.max	Maximum number of iterations. Default value is 500.
stop	Convergence threshold of relative change in cluster means. Default value is 1e-3.
trueY	A vector of true cluster labels of each observation. Default value is NULL.

Details

The `tune_u_joint` function searches over all the combinations of $u \equiv (u_1, \dots, u_M)$ in the neighborhood of \tilde{u} , $\mathcal{N}(\tilde{u}) = \{u : \max(1, \tilde{u}_m - 2) \leq u_m \leq \min(\tilde{u}_m + 2, p_m), m = 1, \dots, M\}$, that minimizes

$$\text{BIC}(u) = -2 \sum_{i=1}^n \log\left(\sum_{k=1}^K \hat{\pi}_k^u f_k(\mathbf{X}_i; \hat{\boldsymbol{\theta}}^u)\right) + \log(n) \cdot K_u.$$

In the above BIC, $K_u = (K - 1) \prod_{m=1}^M u_m + \sum_{m=1}^M p_m(p_m + 1)/2$ is the total number of parameters in TEMM, $\hat{\pi}_k^u$ and $\hat{\boldsymbol{\theta}}^u$ are the estimated parameters with envelope dimension fixed at u . The `tune_u_joint` function intrinsically selects the initial point and return the optimal estimated labels.

Value

<code>opt.u</code>	Optimal envelope dimension selected.
<code>opt.id</code>	Estimated labels fitted by TEMM with the optimal envelope dimension.
<code>opt.mu</code>	Estimated cluster means fitted by TEMM with the optimal envelope dimension.
<code>bic</code>	BIC value.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

Deng, K. and Zhang, X. (2021). Tensor Envelope Mixture Model for Simultaneous Clustering and Multiway Dimension Reduction. *Biometrics*.

See Also

[TEMM](#), [tune_u_sep](#)

Examples

```
A = array(c(rep(1,20),rep(2,20))+rnorm(40),dim=c(2,2,10))
mytune = tune_u_joint(u_candi=list(1:2,1:2),K=2,A)
```

tune_u_sep

Tuning envelope dimension separately by BIC in TEMM.

Description

Tuning envelope dimension separately by BIC in TEMM.

Usage

```
tune_u_sep(m, u_candi, K, X, C = 1, oneD = TRUE,
iter.max = 500, stop = 0.001, trueY = NULL)
```

Arguments

m	The tensor mode to be tuned, can take value in $1, \dots, M$.
u_candi	A vector of candidate envelope dimension.
K	Number of clusters, greater than or equal to 2.
X	The tensor for clustering, should be array type, the last dimension is the sample size n .
C	Constant in separate BIC criterion. Default value is 1.
oneD	Whether to apply 1D-BIC tuning. Default value is TRUE.
iter.max	Maximum number of iterations. Default value is 500.
stop	Convergence threshold of relative change in cluster means. Default value is $1e-3$.
trueY	A vector of true cluster labels of each observation. Default value is NULL.

Details

For tensor mode $m = 1, \dots, M$, the `tune_u_sep` function selects the envelope dimension \tilde{u}_m by minimizing the following BIC-type criterion over the set $\{0, 1, \dots, p_m\}$,

$$\text{BIC}_m(u_m) = \log |\mathbf{\Gamma}_m^T \widehat{\mathbf{M}}_m \mathbf{\Gamma}_m| + \log |\mathbf{\Gamma}_m^T \widehat{\mathbf{N}}_m^{-1} \mathbf{\Gamma}_m| + C \cdot u_m \log(n)/n.$$

This separate selection over each mode m is less sensitive to the complex interrelationships of each mode of the tensor. The default constant C is set as 1 as suggested by Zhang and Mai (2018).

Value

opt.u	Optimal envelope dimension selected.
bic	BIC value.

Author(s)

Kai Deng, Yuqing Pan, Xin Zhang and Qing Mai

References

- Deng, K. and Zhang, X. (2021). Tensor Envelope Mixture Model for Simultaneous Clustering and Multiway Dimension Reduction. *Biometrics*.
- Zhang, X. and Mai, Q. (2018). Model-free envelope dimension selection. *Electronic Journal of Statistics* 12, 2193-2216.

See Also

[TEMM](#), [tune_u_joint](#)

Examples

```
A = array(c(rep(1,20),rep(2,20))+rnorm(40),dim=c(2,2,10))  
mytune = tune_u_sep(1,1:2,K=2,A)
```

Index

DEEM, [2](#), [3](#), [8–10](#)

TEMM, [4](#), [5](#), [7](#), [12](#), [13](#)

TGMM, [6](#), [6](#), [7](#)

tune_K, [3](#), [8](#), [8](#), [10](#)

tune_lamb, [3](#), [8](#), [9](#), [9](#), [10](#)

tune_u_joint, [6](#), [11](#), [12](#), [13](#)

tune_u_sep, [6](#), [12](#), [12](#), [13](#)