

Package ‘TRES’

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Description Provides three estimators for tensor response regression (TRR) and tensor predictor regression (TPR) models with tensor envelope structure. The three types of estimation approaches are generic and can be applied to any envelope estimation problems. The full Grassmannian (FG) optimization is often associated with likelihood-based estimation but requires heavy computation and good initialization; the one-directional optimization approaches (1D and ECD algorithms) are faster, stable and does not require carefully chosen initial values; the SIMPLS-type is motivated by the partial least squares regression and is computationally the least expensive. For details of TRR, see Li L, Zhang X (2017) <[doi:10.1080/01621459.2016.1193022](https://doi.org/10.1080/01621459.2016.1193022)>. For details of TPR, see Zhang X, Li L (2017) <[doi:10.1080/00401706.2016.1272495](https://doi.org/10.1080/00401706.2016.1272495)>. For details of 1D algorithm, see Cook RD, Zhang X (2016) <[doi:10.1080/10618600.2015.1029577](https://doi.org/10.1080/10618600.2015.1029577)>. For details of ECD algorithm, see Cook RD, Zhang X (2018) <[doi:10.5705/ss.202016.0037](https://doi.org/10.5705/ss.202016.0037)>. For more details of the package, see Zeng J, Wang W, Zhang X (2021) <[doi:10.18637/jss.v099.i12](https://doi.org/10.18637/jss.v099.i12)>.

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Description

Provides the ordinary least squares estimator and the three types of tensor envelope structured estimators for tensor response regression (TRR) and tensor predictor regression (TPR) models. The three types of tensor envelope structured approaches are generic and can be applied to any envelope estimation problems. The full Grassmannian (FG) optimization is often associated with likelihood-based estimation but requires heavy computation and good initialization; the one-directional optimization approaches (1D and ECD algorithms) are faster, stable and does not require carefully chosen initial values; the SIMPLS-type is motivated by the partial least squares regression and is computationally the least expensive.

Author(s)

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References

- Zeng J., Wang W., Zhang X. (2021) TRES: An R Package for Tensor Regression and Envelope Algorithms. *Journal of Statistical Software*, 99(12), 1-31. doi:10.18637/jss.v099.i12.
- Cook, R.D. and Zhang, X. (2016). Algorithms for envelope estimation. *Journal of Computational and Graphical Statistics*, 25(1), pp.284-300.
- Li, L. and Zhang, X. (2017). Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.
- Zhang, X. and Li, L. (2017). Tensor envelope partial least-squares regression. *Technometrics*, 59(4), pp.426-436.
- Cook, R.D. and Zhang, X. (2018). Fast envelope algorithms. *Statistica Sinica*, 28(3), pp.1179-1197.

See Also

Useful links:

- <https://github.com/leozeng15/TRES>
- Report bugs at <https://github.com/leozeng15/TRES/issues>

Examples

```
library(TRES)
## Load data "bat"
data("bat")
x <- bat$x
y <- bat$y

## 1. Fitting with OLS method.
fit_ols <- TRR.fit(x, y, method="standard")
```

```

## Print coefficient
coef(fit_ols)

## Print the summary
summary(fit_ols)

## Extract the mean squared error, p-value and standard error from summary
summary(fit_ols)$mse
summary(fit_ols)$p_val
summary(fit_ols)$se

## Make the prediction on the original dataset
predict(fit_ols, x)

## Draw the plots of two-way coefficient tensor (i.e., matrix) and p-value tensor.
plot(fit_ols)

## 2. Fitting with 1D envelope algorithm. (time-consuming)

fit_1D <- TRR.fit(x, y, u = c(14,14), method="1D") # pass envelope rank (14,14)
coef(fit_1D)
summary(fit_1D)
predict(fit_1D, x)
plot(fit_1D)

```

bat

Bat simulated data

Description

Synthetic data generated from tensor response regression (TRR) model. Each response observation is a two-dimensional image, and each binary predictor observation takes values 0 and 1, representing two groups.

Usage

```
data("bat")
```

Format

A list consisting of four components:

x A 1×20 matrix, each entry takes values 0 and 1, representing two groups.

y A $64 \times 64 \times 20$ tensor, each matrix $y@data[, , i]$ represents an image.

coefficients A $64 \times 64 \times 1$ tensor with the bat pattern.

Gamma A list consisting of two 64×14 envelope basis.

Details

The dataset is generated from the tensor response regression (TRR) model:

$$Y_i = BX_i + \epsilon_i, i = 1, \dots, n,$$

where $n = 20$ and the regression coefficient $B \in R^{64 \times 64}$ is a given image with rank 14, representing the mean difference of the response Y between two groups. To make the model conform to the envelope structure, we construct the envelope basis Γ_k and the covariance matrices $\Sigma_k, k = 1, 2$, of error term as following. With the singular value decomposition of B , namely $B = \Gamma_1 \Lambda \Gamma_2^T$, we choose the envelope basis as $\Gamma_k \in R^{64 \times 14}, k = 1, 2$. Then the envelope dimensions are $u_1 = u_2 = 14$. We generate another two matrices $\Omega_k \in R^{14 \times 14} = A_k A_k^T$ and $\Omega_{0k} \in R^{50 \times 50} = A_{0k} A_{0k}^T$, where $A_k \in R^{14 \times 14}$ and $A_{0k} \in R^{50 \times 50}$ are randomly generated from Uniform(0,1) elementwise. Then we set the covariance matrices $\Sigma_k = \Gamma_k \Omega_k \Gamma_k^T + \Gamma_{0k} \Omega_{0k} \Gamma_{0k}^T$, followed by normalization with their Frobenius norms. We set the first 10 predictors $X_i, i = 1, \dots, 10$, as 1 and the rest as 0. The error term is then generated from two-way tensor (matrix) normal distribution $TN(0; \Sigma_1, \Sigma_2)$.

References

Li, L. and Zhang, X., 2017. Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.

Examples

```
## Fit bat dataset with the tensor response regression model
data("bat")
x <- bat$x
y <- bat$y
# Model fitting with ordinary least square.
fit_std <- TRR.fit(x, y, method="standard")
# Draw the coefficient and p-value plots
plot(fit_std)
```

ECD

ECD algorithm for estimating the envelope subspace

Description

Estimate the envelope subspace with specified dimension based on ECD algorithm proposed by Cook, R. D., & Zhang, X. (2018).

Usage

ECD(M, U, u, ...)

Arguments

- M** The p -by- p positive definite matrix M in the envelope objective function.
- U** The p -by- p positive semi-definite matrix U in the envelope objective function.
- u** An integer between 0 and n representing the envelope dimension.
- ...** Additional user-defined arguments:
- **maxiter**: The maximal number of iterations.
 - **tol**: The tolerance used to assess convergence. See the ECD algorithm in Cook, R. D., & Zhang, X. (2018).
- The default values are: `maxiter=500`; `tol=1e-08`.

Details

Estimate M-envelope of $\text{span}(U)$. The dimension of the envelope is u .

See [FGfun](#) for the generic objective function.

The ECD algorithm is similar to 1D algorithm proposed by Cook, R. D., & Zhang, X. (2016). A fast and stable algorithm is used for solving each individual objective function.

Value

Return the orthogonal basis of the envelope subspace with each column represent the sequential direction. For example, the first column is the most informative direction.

References

Cook, R.D. and Zhang, X., 2018. Fast envelope algorithms. *Statistica Sinica*, 28(3), pp.1179-1197.

Examples

```
##simulate two matrices M and U with an envelope structure#
data <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_ECD <- ECD(M, U, u=5)
subspace(Gamma_ECD, G)
```

Description

EEG images data of subjects in alcoholic and control groups.

Usage

```
data("EEG")
```

Format

A list consisting of two components:

x A binary vector with length of 61.

y A $64 \times 64 \times 61$ tensor, consisting of 61 *channels* by *time* EEG images.

Details

The original EEG data contains 77 alcoholic individuals and 45 controls. To reduce the size, we randomly select 61 samples and obtain 39 alcoholic individuals and 22 controls. Each individual was measured with 64 electrodes placed on the scalp sampled at 256 Hz for 1 sec, resulting an EEG image of 64 channels by 256 time points. More information about data collection and some analysis can be found in Zhang et al. (1995) and Li, Kim, and Altman (2010). To facilitate the analysis, the data is downsized along the time domain by averaging every four consecutive time points, yielding a 64×64 matrix response.

References

URL: <https://archive.ics.uci.edu/ml/datasets/EEG+Database>.

Li, L. and Zhang, X., 2017. Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.

Zhang, X.L., Begleiter, H., Porjesz, B., Wang, W. and Litke, A., 1995. Event related potentials during object recognition tasks. *Brain research bulletin*, 38(6), pp.531-538.

Li, B., Kim, M.K. and Altman, N., 2010. On dimension folding of matrix-or array-valued statistical objects. *The Annals of Statistics*, 38(2), pp.1094-1121.

Examples

```
data("EEG")
x <- EEG$x
y <- EEG$y
## Estimate the envelope dimension, the output should be c(1,1).

u <- TRRdim(x, y)$u
u <- c(1,1)

## Fit the dataset with TRR.fit and draw the coefficient plot and p-value plot
fit_1D <- TRR.fit(x, y, u, method = "1D")
plot(fit_1D, xlab = "Time", ylab = "Channels")

## Uncomment display the plots from different methods.
# fit_ols <- TRR.fit(x, y, method = "standard")
# fit_pls <- TRR.fit(x, y, u, method = "PLS")
# plot(fit_ols, xlab = "Time", ylab = "Channels")
# plot(fit_pls, xlab = "Time", ylab = "Channels")
```

 FGfun

The Objective function and its gradient

Description

Calculates the objective function and its gradient for estimating the M -envelope of $\text{span}(U)$, where M is positive definite and U is positive semi-definite.

Usage

```
FGfun(Gamma, M, U)
```

Arguments

Gamma	Γ matrix in the envelope objective function. A p -by- u matrix.
M	The p -by- p positive definite matrix M in the envelope objective function.
U	The p -by- p positive semi-definite matrix U in the envelope objective function.

Details

The generic objective function $F(\Gamma)$ and its gradient $G(\Gamma)$ are listed below for estimating M -envelope of $\text{span}(U)$. For the detailed description, see Cook, R. D., & Zhang, X. (2016).

$$F(\Gamma) = \log |\Gamma^T M \Gamma| + \log |\Gamma^T (M + U)^{-1} \Gamma|$$

$$G(\Gamma) = dF/d\Gamma = 2M\Gamma(\Gamma^T M \Gamma)^{-1} + 2(M + U)^{-1}\Gamma(\Gamma^T (M + U)^{-1} \Gamma)^{-1}$$

Value

F	The value of the objective function at Gamma.
G	The value of the gradient function at Gamma.

References

Cook, R.D. and Zhang, X., 2016. Algorithms for envelope estimation. *Journal of Computational and Graphical Statistics*, 25(1), pp.284-300.

kroncov

*The covariance estimation of tensor normal distribution***Description**

This function provides the MLE of the covariance matrix of tensor normal distribution, where the covariance has a separable Kronecker structure, i.e. $\Sigma = \Sigma_m \otimes \dots \otimes \Sigma_1$. The algorithm is a generalization of the MLE algorithm in Manceur, A. M., & Dutilleul, P. (2013).

Usage

```
kroncov(Tn, tol = 1e-06, maxiter = 10)
```

Arguments

Tn A $p_1 \times \dots \times p_m \times n$ matrix, array or tensor, where n is the sample size.

tol The convergence tolerance with default value 1e-6. The iteration terminates when $\|\Sigma_i^{(t+1)} - \Sigma_i^{(t)}\|_F < \text{tol}$ for some covariance matrix Σ_i .

maxiter The maximal number of iterations. The default value is 10.

Details

The individual component covariance matrices $\Sigma_i, i = 1, \dots, m$ are not identifiable. To overcome the identifiability issue, each matrix Σ_i is normalized at the end of the iteration such that $\|\Sigma_i\|_F = 1$. And an overall normalizing constant λ is extracted so that the overall covariance matrix Σ is defined as

$$\Sigma = \lambda \Sigma_m \otimes \dots \otimes \Sigma_1.$$

If Tn is a $p \times n$ design matrix for a multivariate random variable, then $\text{lambda} = 1$ and S is a length-one list containing the sample covariance matrix.

Value

lambda The normalizing constant.

S A matrix list, consisting of each normalized covariance matrix $\Sigma_1, \dots, \Sigma_m$.

References

Manceur, A.M. and Dutilleul, P., 2013. Maximum likelihood estimation for the tensor normal distribution: Algorithm, minimum sample size, and empirical bias and dispersion. *Journal of Computational and Applied Mathematics*, 239, pp.37-49.

manifold1D

*Estimate the envelope subspace (ManifoldOptim 1D)***Description**

The 1D algorithm (Cook and Zhang 2016) implemented with Riemannian manifold optimization from R package **ManifoldOptim**.

Usage

```
manifold1D(M, U, u, ...)
```

Arguments

- | | |
|-----|--|
| M | The p -by- p positive definite matrix M in the envelope objective function. |
| U | The p -by- p positive semi-definite matrix U in the envelope objective function. |
| u | An integer between 0 and n representing the envelope dimension. |
| ... | Additional user-defined arguments: <ul style="list-style-type: none"> • <code>maxiter</code>: The maximal number of iterations. • <code>tol</code>: The tolerance used to assess convergence. See Huang et al. (2018) for details on how this is used. • <code>method</code>: The name of optimization method supported by R package ManifoldOptim. <ul style="list-style-type: none"> – "LRBFGS": Limited-memory RBFGRS – "LRTRSR1": Limited-memory RTRSR1 – "RBFGRS": Riemannian BFGS – "RBroydenFamily": Riemannian Broyden family – "RCG": Riemannian conjugate gradients – "RNewton": Riemannian line-search Newton – "RSD": Riemannian steepest descent – "RTRNewton": Riemannian trust-region Newton – "RTRSD": Riemannian trust-region steepest descent – "RTRSR1": Riemannian trust-region symmetric rank-one update – "RWRBFGS": Riemannian BFGS • <code>check</code>: Logical value. Should internal manifold object check inputs and print summary message before optimization. |

The default values are: `maxiter = 500`; `tol = 1e-08`; `method = "RCG"`; `check = FALSE`.

Details

Estimate M-envelope of $\text{span}(U)$. The dimension of the envelope is u .

Value

Return the estimated orthogonal basis of the envelope subspace.

References

Cook, R.D. and Zhang, X., 2016. Algorithms for envelope estimation. *Journal of Computational and Graphical Statistics*, 25(1), pp.284-300.

Huang, W., Absil, P.A., Gallivan, K.A. and Hand, P., 2018. ROPTLIB: an object-oriented C++ library for optimization on Riemannian manifolds. *ACM Transactions on Mathematical Software (TOMS)*, 44(4), pp.1-21.

See Also

[MenvU_sim](#), [subspace](#)

Examples

```
## Simulate two matrices M and U with an envelope structure
data <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_1D <- manifold1D(M, U, u = 5)
subspace(Gamma_1D, G)
```

manifoldFG

Estimate the envelope subspace (ManifoldOptim FG)

Description

The FG algorithm (Cook and Zhang 2016) implemented with Riemannian manifold optimization from R package **ManifoldOptim**.

Usage

```
manifoldFG(M, U, u, Gamma_init = NULL, ...)
```

Arguments

M	The p -by- p positive definite matrix M in the envelope objective function.
U	The p -by- p positive semi-definite matrix U in the envelope objective function.
u	An integer between 0 and n representing the envelope dimension. Ignored if <code>Gamma_init</code> is provided.
Gamma_init	Initial envelope subspace basis. The default value is the estimator from <code>manifold1D(M, U, u)</code> .

... Additional user-defined arguments:

- `maxiter`: The maximal number of iterations.
- `tol`: The tolerance used to assess convergence. See Huang et al. (2018) for details on how this is used.
- `method`: The name of optimization method supported by R package **ManifoldOptim**
 - "LRBFGS": Limited-memory RBFSS
 - "LRTRSR1": Limited-memory RTRSR1
 - "RBFSS": Riemannian BFGS
 - "RBroydenFamily": Riemannian Broyden family
 - "RCG": Riemannian conjugate gradients
 - "RNewton": Riemannian line-search Newton
 - "RSD": Riemannian steepest descent
 - "RTRNewton": Riemannian trust-region Newton
 - "RTRSD": Riemannian trust-region steepest descent
 - "RTRSR1": Riemannian trust-region symmetric rank-one update
 - "RWRBFGS": Riemannian BFGS
- `check`: Logical value. Should internal manifold object check inputs and print summary message before optimization.

The default values are: `maxiter = 500`; `tol = 1e-08`; `method = "RCG"`; `check = FALSE`.

Details

Estimate M-envelope of $\text{span}(U)$. The dimension of the envelope is u .

Value

Return the estimated orthogonal basis of the envelope subspace.

References

Cook, R.D. and Zhang, X., 2016. Algorithms for envelope estimation. *Journal of Computational and Graphical Statistics*, 25(1), pp.284-300.

Huang, W., Absil, P.A., Gallivan, K.A. and Hand, P., 2018. ROPTLIB: an object-oriented C++ library for optimization on Riemannian manifolds. *ACM Transactions on Mathematical Software (TOMS)*, 44(4), pp.1-21.

Examples

```
##simulate two matrices M and U with an envelope structure
data <- MenvU_sim(p=20, u=5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_FG <- manifoldFG(M, U, u=5)
subspace(Gamma_FG, G)
```

MenvU_sim	<i>Generate matrices M and U</i>
-----------	----------------------------------

Description

This function generates the matrices M and U with envelope structure.

Usage

```
MenvU_sim(
  p,
  u,
  Omega = NULL,
  Omega0 = NULL,
  Phi = NULL,
  jitter = FALSE,
  wishart = FALSE,
  n = NULL
)
```

Arguments

<code>p</code>	Dimension of p -by- p matrix M .
<code>u</code>	The envelope dimension. An integer between 0 and p .
<code>Omega</code>	The positive definite matrix Ω in $M = \Gamma\Omega\Gamma^T + \Gamma_0\Omega_0\Gamma_0^T$. The default is $\Omega = AA^T$ where the elements in A are generated from Uniform(0,1) distribution.
<code>Omega0</code>	The positive definite matrix Ω_0 in $M = \Gamma\Omega\Gamma^T + \Gamma_0\Omega_0\Gamma_0^T$. The default is $\Omega_0 = AA^T$ where the elements in A are generated from Uniform(0,1) distribution.
<code>Phi</code>	The positive definite matrix Φ in $U = \Gamma\Phi\Gamma^T$. The default is $\Phi = AA^T$ where the elements in A are generated from Uniform(0,1) distribution.
<code>jitter</code>	Logical or numeric. If it is numeric, the diagonal matrix <code>diag(jitter, nrow(M), ncol(M))</code> is added to matrix M to ensure the positive definiteness of M . If it is TRUE, then it is set as $1e-5$ and the jitter is added. If it is FALSE (default), no jitter is added.
<code>wishart</code>	Logical. If it is TRUE, the sample estimator from Wishart distribution $W_p(M/n, n)$ and $W_p(U/n, n)$ are generated as the output matrices M and U .
<code>n</code>	The sample size. If <code>wishart</code> is FALSE, then <code>n</code> is ignored.

Details

The matrices M and U are in forms of

$$M = \Gamma\Omega\Gamma^T + \Gamma_0\Omega_0\Gamma_0^T, U = \Gamma\Phi\Gamma^T.$$

The envelope basis Γ is randomly generated from the Uniform (0, 1) distribution elementwise and then transformed to a semi-orthogonal matrix. Γ_0 is the orthogonal completion of Γ .

In some cases, to guarantee that M is positive definite which is required by the definition of envelope, a jitter should be added to M .

If `wishart` is TRUE, after the matrices M and U are generated, the samples from Wishart distribution $W_p(M/n, n)$ and $W_p(U/n, n)$ are output as matrices M and U . If so, n is required.

Value

M	The p -by- p matrix M .
U	The p -by- p matrix U .
Gamma	The p -by- u envelope basis.

References

Cook, R.D. and Zhang, X., 2018. Fast envelope algorithms. *Statistica Sinica*, 28(3), pp.1179-1197.

Examples

```
data1 <- MenvU_sim(p = 20, u = 5)
M1 <- data1$M
U1 <- data1$U

# Sample version from Wishart distribution
data2 <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M2 <- data2$M
U2 <- data2$U
```

oneD_bic

Envelope dimension selection based on 1D-BIC

Description

This function selects envelope subspace dimension using 1D-BIC proposed by Zhang, X., & Mai, Q. (2018). The constrained optimization in the 1D algorithm is based on the line search algorithm for optimization on manifold. The algorithm is developed by Wen and Yin (2013) and the Matlab version is in the Matlab package **OptM**.

Usage

```
oneD_bic(M, U, n, C = 1, maxdim = 10, ...)
```

Arguments

M	The p -by- p positive definite matrix M in the envelope objective function.
U	The p -by- p positive semi-definite matrix U in the envelope objective function.
n	The sample size.
C	The constant defined in 1D-BIC criterion, the default value is 1.

maxdim	The maximum dimension to consider, maxdim is smaller than p , the default value is 10.
...	Additional user-defined arguments for the line search algorithm: <ul style="list-style-type: none"> • maxiter: The maximal number of iterations. • xtol: The convergence tolerance for the relative changes of the consecutive iterates w, e.g., $\ w^{(k)} - w^{(k-1)}\ _F / \sqrt{p}$ • gtol: The convergence tolerance for the gradient of Lagrangian, e.g., $\ G^{(k)} - w^{(k)}(G^{(t)})^T w^{(t)}\ _F$ • ftol: The convergence tolerance for relative changes of the consecutive objective function values F, e.g., $F^{(k)} - F^{(k-1)} / (1 + F^{(k-1)})$. Usually, $\max\{xtol, gtol\} > ftol$
	The default values are: maxiter=500; xtol=1e-08; gtol=1e-08; ftol=1e-12.

Details

The objective function $F(w)$ and its gradient $G(w)$ in line search algorithm are:

$$F(w) = \log |w^T M_k w| + \log |w^T (M_k + U_k)^{-1} w|$$

$$G(w) = dF/dw = 2(w^T M_k w)^{-1} M_k w + 2(w^T (M_k + U_k)^{-1} w)^{-1} (M_k + U_k)^{-1} w$$

See Cook, R. D., & Zhang, X. (2016) for more details of the 1D algorithm.

The 1D-BIC criterion is defined as

$$I(k) = \sum_{j=1}^k \phi_j(\hat{w}_j) + Ck \log(n)/n, \quad k = 0, 1, \dots, p,$$

where $C > 0$ is a constant, \hat{w} is the 1D solver, the function ϕ_j is the individual objective function solved by 1D algorithm, n is the sample size. Then the selected dimension u is the one yielding the smallest 1D-BIC $I(k)$. See Zhang, X., & Mai, Q. (2018) for more details.

As suggested by Zhang, X., & Mai, Q. (2018), the number C should be set to its default value $C = 1$ when there is no additional model assumption or prior information. However, if additional model assumption or prior information are known, C should be set such that Ck best matches the degree-of-freedom or total number of free parameters of the model or estimation procedure. For example, in TRR model where the predictor design matrix is of dimension $p \times n$, C should be set as p . See Zhang, X., & Mai, Q. (2018) for more details.

Value

bicval	The BIC values for different envelope dimensions.
u	The dimension selected which corresponds to the smallest BIC values.
Gamma	The estimation of envelope subspace basis.

References

- Zhang, X. and Mai, Q., 2018. Model-free envelope dimension selection. *Electronic Journal of Statistics*, 12(2), pp.2193-2216.
- Wen, Z. and Yin, W., 2013. A feasible method for optimization with orthogonality constraints. *Mathematical Programming*, 142(1-2), pp.397-434.

See Also

[OptM1D](#), [MenvU_sim](#)

Examples

```
##simulate two matrices M and U with an envelope structure
data <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
bic <- oneD_bic(M, U, n = 200)
## visualization
plot(1:10, bic$bicval, type="o", xlab="Envelope Dimension", ylab="BIC values",
main="Envelope Dimension Selection")
```

OptM1D

Estimate the envelope subspace (OptM 1D)

Description

The 1D algorithm to estimate the envelope subspace based on the line search algorithm for optimization on manifold. The line search algorithm is developed by Wen and Yin (2013) and the Matlab version is implemented in the Matlab package **OptM**.

Usage

OptM1D(M, U, u, ...)

Arguments

- | | |
|-----|--|
| M | The p -by- p positive definite matrix M in the envelope objective function. |
| U | The p -by- p positive semi-definite matrix U in the envelope objective function. |
| u | An integer between 0 and n representing the envelope dimension. |
| ... | Additional user-defined arguments for the line search algorithm: <ul style="list-style-type: none"> • <code>maxiter</code>: The maximal number of iterations. • <code>xtol</code>: The convergence tolerance for the relative changes of the consecutive iterates w, e.g., $\ w^{(k)} - w^{(k-1)}\ _F / \sqrt{p}$ • <code>gtol</code>: The convergence tolerance for the gradient of Lagrangian, e.g., $\ G^{(k)} - w^{(k)}(G^{(k)})^T w^{(k)}\ _F$ • <code>ftol</code>: The convergence tolerance for relative changes of the consecutive objective function values F, e.g., $F^{(k)} - F^{(k-1)} / (1 + F^{(k-1)})$. Usually, $\max\{xtol, gtol\} > ftol$ |

The default values are: `maxiter=500`; `xtol=1e-08`; `gtol=1e-08`; `ftol=1e-12`.

Details

The objective function $F(w)$ and its gradient $G(w)$ in line search algorithm are:

$$F(w) = \log |w^T M_k w| + \log |w^T (M_k + U_k)^{-1} w|$$

$$G(w) = dF/dw = 2(w^T M_k w)^{-1} M_k w + 2(w^T (M_k + U_k)^{-1} w)^{-1} (M_k + U_k)^{-1} w$$

See Cook, R. D., & Zhang, X. (2016) for more details of the 1D algorithm.

Value

Return the estimated orthogonal basis of the envelope subspace.

References

Cook, R.D. and Zhang, X., 2016. Algorithms for envelope estimation. *Journal of Computational and Graphical Statistics*, 25(1), pp.284-300.

Wen, Z. and Yin, W., 2013. A feasible method for optimization with orthogonality constraints. *Mathematical Programming*, 142(1-2), pp.397-434.

Examples

```
## Simulate two matrices M and U with an envelope structure
data <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_1D <- OptM1D(M, U, u = 5)
subspace(Gamma_1D, G)
```

OptMFG

Estimate the envelope subspace (OptM FG)

Description

The FG algorithm to estimate the envelope subspace based on the curvilinear search algorithm for optimization on Stiefel manifold. The curvilinear algorithm is developed by Wen and Yin (2013) and the Matlab version is implemented in the Matlab package **OptM**.

Usage

```
OptMFG(M, U, u, Gamma_init = NULL, ...)
```

Arguments

<code>M</code>	The p -by- p positive definite matrix M in the envelope objective function.
<code>U</code>	The p -by- p positive semi-definite matrix U in the envelope objective function.
<code>u</code>	An integer between 0 and n representing the envelope dimension. Ignored if <code>Gamma_init</code> is provided.
<code>Gamma_init</code>	Initial envelope subspace basis. The default value is the estimator from <code>OptM1D(M, U, u)</code> .
<code>...</code>	Additional user-defined arguments for the curvilinear search algorithm: <ul style="list-style-type: none"> <code>maxiter</code>: The maximal number of iterations. <code>xtol</code>: The convergence tolerance for Γ, e.g., $\ \Gamma^{(k)} - \Gamma^{(k-1)}\ _F / \sqrt{p}$ <code>gtol</code>: The convergence tolerance for the projected gradient, e.g., $\ G^{(k)} - \Gamma^{(k)}(G^{(k)})^T \Gamma^{(k)}\ _F$ <code>ftol</code>: The convergence tolerance for objective function F, e.g., $F^{(k)} - F^{(k-1)} / (1 + F^{(k-1)})$. Usually, $\max\{xtol, gtol\} > ftol$ The default values are: <code>maxiter=500</code> ; <code>xtol=1e-08</code> ; <code>gtol=1e-08</code> ; <code>ftol=1e-12</code> .

Details

If `Gamma_init` is provided, then the envelope dimension `u = ncol(Gamma_init)`.

The function `OptMFG` calls the function `OptStiefelGGB` internally which implements the curvilinear search algorithm.

The objective function $F(\Gamma)$ and its gradient $G(\Gamma)$ in curvilinear search algorithm are:

$$F(\Gamma) = \log |\Gamma^T M \Gamma| + \log |\Gamma^T (M + U)^{-1} \Gamma|$$

$$G(\Gamma) = dF/d\Gamma = 2M\Gamma(\Gamma^T M \Gamma)^{-1} + 2(M + U)^{-1}\Gamma(\Gamma^T (M + U)^{-1} \Gamma)^{-1}$$

Value

Return the estimated orthogonal basis of the envelope subspace.

References

Wen, Z. and Yin, W., 2013. A feasible method for optimization with orthogonality constraints. *Mathematical Programming*, 142(1-2), pp.397-434.

See Also

[OptStiefelGGB](#)

Examples

```
##simulate two matrices M and U with an envelope structure
data <- MenvU_sim(p=20, u=5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_FG <- OptMFG(M, U, u=5)
subspace(Gamma_FG, G)
```

OptStiefelGBB	<i>Optimization on Stiefel manifold</i>
---------------	---

Description

Curvilinear search algorithm for optimization on Stiefel manifold developed by Wen and Yin (2013).

Usage

OptStiefelGBB(X, fun, opts = NULL, ...)

Arguments

X	Initial value to start the optimization. A n by k orthonormal matrix such that $X^T X = I_k$.
fun	The function that returns the objective function value and its gradient. The syntax for fun is fun(X, data1, data2) where data1, data2 are additional data passed to ...
opts	A list specifying additional user-defined arguments for the curvilinear search algorithm. Some important ones are listed in the following: <ul style="list-style-type: none"> • maxiter: The maximal number of iterations. • xtol: The convergence tolerance for X, e.g., $\ X^{(t)} - X^{(t-1)}\ _F / \sqrt{k}$. • gtol: The convergence tolerance for the gradient of the Lagrangian function, e.g., $\ G^{(t)} - X^{(t)}(G^{(t)})^T X^{(t)}\ _F$. • ftol: The convergence tolerance for objective function F, e.g., $F^{(t)} - F^{(t-1)} / (1 + F^{(t-1)})$. Usually, $\max\{xtol, gtol\} > ftol$. <p>The default values are: maxiter=500; xtol=1e-08; gtol=1e-08; ftol=1e-12.</p>
...	Additional input passed to fun.

Details

The calling syntax is OptStiefelGBB(X, fun, opts, data1, data2), where fun(X, data1, data2) returns the objective function value and its gradient.

For example, for n by k matrix X , the optimization problem is

$$\min_X -tr(X^T W X), \text{ such that } X^T X = I_k.$$

The objective function and its gradient are

$$F(X) = -tr(X^T W X), G(X) = -2W X.$$

Then we need to provide the function fun(X, W) which returns $F(X)$ and $G(X)$. See **Examples** for details.

For more details of the termination rules and the tolerances, we refer the interested readers to Section 5.1 of Wen and Yin (2013).

Value

<code>X</code>	The converged solution of the optimization problem.
<code>out</code>	Output information, including estimation error, function value, iteration times etc. <ul style="list-style-type: none"> • <code>nfe</code>: The total number of line search attempts. • <code>msg</code>: Message: "convergence" "exceed max iteration". • <code>feasi</code>: The feasibility of solution: $\ X^T X - I_k\ _F$. • <code>nrmG</code>: The convergence criterion based on the projected gradient $\ G - XG^T X\ _F$. • <code>fval</code>: The objective function value $F(X)$ at termination. • <code>iter</code>: The number of iterations.

References

Wen, Z. and Yin, W., 2013. A feasible method for optimization with orthogonality constraints. *Mathematical Programming*, 142(1-2), pp.397-434.

Examples

```
n <- 1000
k <- 6

# Randomly generated matrix M
W <- matrix(rnorm(n^2), n, n)
W <- t(W) %*% W

# Randomly generated orthonormal initial matrix
X0 <- matrix(rnorm(n*k), n, k)
X0 <- qr.Q(qr(X0))

# The objective function and its gradient
fun <- function(X, W){
  F <- - sum(diag(t(X) %*% W %*% X))
  G <- - 2*(W %*% X)
  return(list(F = F, G = G))
}

# Options list
opts<-list(record = 0, maxiter = 1000, xtol = 1e-5, gtol = 1e-5, ftol = 1e-8)

# Main part
output <- OptStiefelGGB(X0, fun, opts, W)
X <- output$X
out <- output$out
```

plot.Tenv *Plot coefficients and p-value for Tenv object.*

Description

Plot method for object returned from TRR.fit and TPR.fit functions.

Usage

```
## S3 method for class 'Tenv'
plot(
  x,
  level = 0.05,
  main = paste0("Coefficient plot ", "(" , x$method, ")"),
  main_p = paste0("P value plot ", "(" , x$method, ")"),
  xlab = "",
  ylab = "",
  axes = TRUE,
  ask = TRUE,
  ...
)
```

Arguments

x	An object of class "Tenv", as the ones returned from TPR.fit or TRR.fit.
level	The significant level of p-value. Default is 0.05.
main	The title of coefficient plot.
main_p	The title of p-value plot.
xlab	The title of x-axis.
ylab	The title of y-axis.
axes	A logical value specifying whether the axes should be drawn.
ask	A logical value. If it is TRUE (default), user is prompted before the second plot is shown (if exists).
...	Other parameters to be passed to the plotting functions.

Details

coef(x) must be a two-way tensor or a matrix.

Since p -value depend on $\widehat{\text{cov}}^{-1}\{\text{vec}(\mathbf{X})\}$ which is unavailable for the ultra-high dimensional $\text{vec}(\mathbf{X})$ in tensor predictor regression (TPR), the p -value plot is not provided for the object returned from TPR.fit. Therefore, for the object return from TPR.fit, only the coefficients plot is displayed. And for the object return from TRR.fit, both the coefficients plot and p -value plot are displayed.

main and main_p control the titles of coefficient plot and p -value plot separately. Some other arguments used in function graphics::image, e.g., xlim, ylim, zlim, col, xaxs, yaxs, etc., can be passed to ...

ask can be set as FALSE if the pause before the second plot is not preferred. If x is an object from TPR.fit, no pause is enabled.

Value

No return value.

See Also

[TRR.fit](#), [TPR.fit](#)

Examples

```
data("bat")
x <- bat$x
y <- bat$y
fit <- TRR.fit(x, y, method="standard")
plot(fit)

## Change the significant level to 0.1
plot(fit, level = 0.1)
```

PMSE

Prediction and mean squared error.

Description

Evaluate the tensor response regression (TRR) or tensor predictor regression (TPR) model through the mean squared error.

Usage

```
PMSE(x, y, B)
```

Arguments

x	A predictor tensor, array, matrix or vector.
y	A response tensor, array, matrix or vector.
B	An coefficient tensor tensor, array, matrix or vector.

Details

There are three situations:

- TRR model: If y is an m -way tensor (array), x should be matrix or vector and B should be tensor or array.
- TPR model: If x is an m -way tensor (array), y should be matrix or vector and B should be tensor or array.

- Other: If x and y are both matrix or vector, B should be matrix. In this case, the prediction is calculated as $\text{pred} = B \cdot X$.

In any cases, users are asked to ensure the dimensions of x , y and B match. See [TRRsim](#) and [TPRsim](#) for more details of the TRR and TPR models.

Let \hat{Y}_i denote each prediction, then the mean squared error is defined as $1/n \sum_{i=1}^n \|Y_i - \hat{Y}_i\|_F^2$, where $\|\cdot\|_F$ denotes the Frobenius norm.

Value

mse	The mean squared error.
pred	The predictions.

See Also

[TRRsim](#), [TPRsim](#).

Examples

```
## Dataset in TRR model
r <- c(10, 10, 10)
u <- c(2, 2, 2)
p <- 5
n <- 100
dat <- TRRsim(r = r, p = p, u = u, n = n)
x <- dat$x
y <- dat$y

# Fit data with TRR.fit
fit_std <- TRR.fit(x, y, method="standard")
result <- PMSE(x, y, fit_std$coefficients)
## Dataset in TPR model
p <- c(10, 10, 10)
u <- c(1, 1, 1)
r <- 5
n <- 200
dat <- TPRsim(p = p, r = r, u = u, n = n)
x <- dat$x
y <- dat$y

# Fit data with TPR.fit
fit_std <- TPR.fit(x, y, u, method="standard")
result <- PMSE(x, y, fit_std$coefficients)
```

predict.Tenv	<i>Predict method for Tenv object.</i>
--------------	--

Description

Predict response for the object returned from `TRR.fit` and `TPR.fit` functions.

Usage

```
## S3 method for class 'Tenv'
predict(object, newdata, ...)
```

Arguments

object	An object of class "Tenv", as the ones returned from <code>TPR.fit</code> or <code>TRR.fit</code> .
newdata	The data to be used for prediction. It can be a vector, a matrix or a tensor if object is returned from <code>TRR.fit</code> , and can be a matrix or a tensor if object is returned from <code>TPR.fit</code> .
...	Additional arguments. No available arguments exist in this version.

Value

Return the predicted response.

Note

If newdata is missing, the fitted response from object is returned.

Examples

```
data("bat")
x <- bat$x
y <- bat$y
fit <- TRR.fit(x, y, method="standard")
predict(fit, x)
```

simplsMU	<i>SIMPLS-type algorithm for estimating the envelope subspace</i>
----------	---

Description

This algorithm is a generalization of the SIMPLS algorithm in De Jong, S. (1993). See Cook (2018) Section 6.5 for more details of this generalized moment-based envelope algorithm; see Cook, Helland, and Su (2013) for a connection between SIMPLS and the predictor envelope in linear model.

Usage

```
simplsMU(M, U, u)
```

Arguments

M The p -by- p positive definite matrix M in the envelope objective function.

U The p -by- p positive semi-definite matrix U in the envelope objective function.

u An integer between 0 and n representing the envelope dimension.

Value

Returns the estimated orthogonal basis of the envelope subspace.

References

De Jong, S., 1993. SIMPLS: an alternative approach to partial least squares regression. *Chemometrics and intelligent laboratory systems*, 18(3), pp.251-263.

Cook, R.D., Helland, I.S. and Su, Z., 2013. Envelopes and partial least squares regression. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, 75(5), pp.851-877.

Cook, R.D., 2018. An introduction to envelopes: dimension reduction for efficient estimation in multivariate statistics (Vol. 401). John Wiley & Sons.

Examples

```
##simulate two matrices M and U with an envelope structure#
data <- MenvU_sim(p = 20, u = 5, wishart = TRUE, n = 200)
M <- data$M
U <- data$U
G <- data$Gamma
Gamma_pls <- simplsMU(M, U, u=5)
subspace(Gamma_pls, G)
```

square

Square simulated data

Description

Synthetic data generated from tensor predictor regression (TPR) model. Each response observation is univariate, and each predictor observation is a matrix.

Usage

```
data("square")
```

Format

A list consisting of four components:

x A $32 \times 32 \times 200$ tensor, each matrix `x@data[, , i]` represents a predictor observation.

y A 1×200 matrix, each entry represents a response observation.

coefficients A $32 \times 32 \times 1$ tensor with a square pattern.

Gamma A list consisting of two 32×2 envelope basis.

Details

The dataset is generated from the tensor predictor regression (TPR) model:

$$Y_i = B_{(m+1)} \text{vec}(X_i) + \epsilon_i, \quad i = 1, \dots, n,$$

where $n = 200$ and the regression coefficient $B \in R^{32 \times 32}$ is a given image with rank 2, which has a square pattern. All the elements of the coefficient matrix B are either 0.1 or 1. To make the model conform to the envelope structure, we construct the envelope basis Γ_k and the covariance matrices $\Sigma_k, k = 1, 2$, of predictor X as following. With the singular value decomposition of B , namely $B = \Gamma_1 \Lambda \Gamma_2^T$, we choose the envelope basis as $\Gamma_k \in R^{32 \times 2}, k = 1, 2$. Then the envelope dimensions are $u_1 = u_2 = 2$. We set matrices $\Omega_k = I_2$ and $\Omega_{0k} = 0.01 I_{30}, k = 1, 2$. Then we generate the covariance matrices $\Sigma_k = \Gamma_k \Omega_k \Gamma_k^T + \Gamma_{0k} \Omega_{0k} \Gamma_{0k}^T$, followed by normalization with their Frobenius norms. The predictor X_i is then generated from two-way tensor (matrix) normal distribution $TN(0; \Sigma_1, \Sigma_2)$. And the error term ϵ_i is generated from standard normal distribution.

References

Zhang, X. and Li, L., 2017. Tensor envelope partial least-squares regression. *Technometrics*, 59(4), pp.426-436.

Examples

```
## Fit square dataset with the tensor predictor regression model
data("square")
x <- square$x
y <- square$y
# Model fitting with ordinary least square.
fit_std <- TPR.fit(x, y, method="standard")
# Draw the coefficient plot.
plot(fit_std)
```

std_err	<i>Elementwise standard error.</i>
---------	------------------------------------

Description

Calculates the elementwise standard error for the object returned from `TRR.fit`. The standard error for the object returned from `TPR.fit` is unavailable.

Usage

```
std_err(object)
```

Arguments

`object` an object of class "Tenv", as the ones returned from `TRR.fit`.

Value

The standard error tensor is returned.

Note

The function only supports the object returned from `TRR.fit` since there is no standard error for the object returned from `TPR.fit`.

Examples

```
data("bat")
x <- bat$x
y <- bat$y
fit <- TRR.fit(x, y, method="standard")
std_err(fit)
```

subspace	<i>The distance between two subspaces.</i>
----------	--

Description

This function calculates the distance between the two subspaces with equal dimensions $\text{span}(A)$ and $\text{span}(B)$, where $A \in R^{p \times u}$ and $B \in R^{p \times u}$ are the basis matrices of two subspaces. The distance is defined as

$$\|P_A - P_B\|_F / \sqrt{2d},$$

where P is the projection matrix onto the given subspace with the standard inner product, and d is the common dimension.

Usage

```
subspace(A, B)
```

Arguments

A A p -by- u full column rank matrix.
 B A p -by- u full column rank matrix.

Value

Returns a distance metric that is between 0 and 1

summary.Tenv	<i>Summarize method for Tenv object.</i>
--------------	--

Description

Summary method for object returned from `TRR.fit` and `TPR.fit` functions.

Usage

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

## S3 method for class 'summary.Tenv'
print(x, ...)
```

Arguments

object An object of class "Tenv", as the ones returned from `TPR.fit` or `TRR.fit`.
 ... Additional arguments. No available arguments exist in this version.
 x An object of class "summary.Tenv", usually, a result of a call to `summary.Tenv`.

Details

Extract call, method, coefficients, residuals, Gamma from object. And append mse, p -value and the standard error of estimated coefficient.

The mean squared error mse is defined as $1/n \sum_{i=1}^n \|Y_i - \hat{Y}_i\|_F^2$, where \hat{Y}_i is the prediction and $\|\cdot\|_F$ is the Frobenius norm of tensor.

Since the p -value and standard error depend on the estimation of $\text{cov}^{-1}(\text{vec}(X))$ which is unavailable for the ultra-high dimensional $\text{vec}(X)$ in tensor predictor regression (TPR), the two statistics are only provided for the object returned from `TRR.fit`.

`print.summary.Tenv` provides a more readable form of the statistics contained in `summary.Tenv`. If object is returned from `TRR.fit`, then `p-val` and `se` are also returned.

Value

Return object with additional components

call	The matched call.
method	The implemented method.
n	The sample size.
xdim	The dimension of predictor.
ydim	The dimension of response.
coefficients	The tensor coefficients estimated from <code>TPR.fit</code> or <code>TRR.fit</code> .
residuals	The residuals, which equals to the response minus the fitted values.
Gamma	A list of envelope subspace basis.
mse	The mean squared error. The mean squared Frobenius norm of the difference between each response Y_i and fitted value \hat{Y}_i .
p_val	The p-value for coefficients. Only for the object returned from <code>TRR.fit</code> .
se	The standard error for coefficients. Only for the object returned from <code>TRR.fit</code> .

See Also

Fitting functions [TRR.fit](#), [TPR.fit](#).

Examples

```
data("bat")
x <- bat$x
y <- bat$y
fit <- TRR.fit(x, y, method="standard")
##print summary
summary(fit)

##Extract the p-value and standard error from summary
summary(fit)$p_val
summary(fit)$se
```

Tenv_Pval

The p-value and standard error of coefficient in tensor response regression (TRR) model.

Description

Obtain p -value of each element in the tensor regression coefficient estimator. Two-sided t-tests are implemented on the coefficient estimator, where asymptotic covariance of the OLS estimator is used.

Usage

```
Tenv_Pval(x, y, Bhat)
```

Arguments

x The response tensor instance $r_1 \times r_2 \times \dots \times r_m$.

y A vector predictor of dimension p .

Bhat The estimator of tensor regression coefficient.

The p -value and the standard error of estimated coefficient are not provided for tensor predictor regression since they depend on $\widehat{\text{cov}}^{-1}\{\text{vec}(\mathbf{X})\}$ which is unavailable due to the ultra-high dimension of $\text{vec}(\mathbf{X})$.

Value

p_ols The p-value tensor of OLS estimator.

p_val The p-value tensor of Bhat.

se The standard error tensor of Bhat.

Examples

```
## Use dataset bat
data("bat")
x <- bat$x
y <- bat$y
fit_std <- TRR.fit(x, y, method="standard")
Tenv_Pval(x, y, fit_std$coefficients)
```

 TPR.fit

Tensor predictor regression

Description

This function is used for estimation of tensor predictor regression. The available method including standard OLS type estimation, PLS type of estimation as well as envelope estimation with FG, 1D and ECD approaches.

Usage

```
TPR.fit(x, y, u, method=c('standard', 'FG', '1D', 'ECD', 'PLS'), Gamma_init = NULL)
```

Arguments

x	The predictor tensor instance of dimension $p_1 \times p_2 \times \dots \times p_m \times n$, where n is the sample size. Array with the same dimensions and matrix with dimension $p \times n$ are acceptable. If y is missing, x should be a list or an environment consisting of predictor and response datasets.
y	The response matrix of dimension $r \times n$, where n is the sample size. Vector of length n is acceptable.
u	The dimension of envelope subspace. $u = (u_1, \dots, u_m)$. Used for methods "FG", "1D", "ECD" and "PLS". User can use TPRdim to select dimension.
method	The method used for estimation of tensor response regression. There are four possible choices. <ul style="list-style-type: none"> • "standard": The standard OLS type estimation. • "FG": Envelope estimation with full Grassmannian (FG) algorithm. • "1D": Envelope estimation with one dimensional optimization approaches by 1D algorithm. • "ECD": Envelope estimation with one dimensional optimization approaches by ECD algorithm. • "PLS": The SIMPLS-type estimation without manifold optimization.
Gamma_init	A list specifying the initial envelope subspace basis for "FG" method. By default, the estimators given by "1D" algorithm is used.

Details

Please refer to **Details** part of [TPRsim](#) for the description of the tensor predictor regression model.

Value

`TPR.fit` returns an object of class "Tenv".

The function [summary](#) (i.e., [summary.Tenv](#)) is used to print the summary of the results, including additional information, e.g., the p-value and the standard error for coefficients, and the prediction mean squared error.

The functions `coefficients`, `fitted.values` and `residuals` can be used to extract different features returned from `TPR.fit`.

The function [plot](#) (i.e., [plot.Tenv](#)) plots the two-dimensional coefficients and p-value for object of class "Tenv".

The function [predict](#) (i.e., [predict.Tenv](#)) predicts response for the object returned from `TPR.fit` function.

x	The original predictor dataset.
y	The original response dataset.
call	The matched call.
method	The implemented method.
coefficients	The estimation of regression coefficient tensor.
Gamma	The estimation of envelope subspace basis.

Sigma	A lists of estimated covariance matrices at each mode for the tensor predictors.
fitted.values	The fitted response matrix.
residuals	The residuals matrix.

References

Zhang, X. and Li, L., 2017. Tensor envelope partial least-squares regression. *Technometrics*, 59(4), pp.426-436.

See Also

[summary.Tenv](#) for summaries, calculating mean squared error from the prediction.
[plot.Tenv](#)(via `graphics::image`) for drawing the two-dimensional coefficient plot.
[predict.Tenv](#) for prediction.
 The generic functions [coef](#), [residuals](#), [fitted](#).
[TPRdim](#) for selecting the dimension of envelope by cross-validation.
[TPRsim](#) for generating the simulated data used in tensor prediction regression.
 The simulated data [square](#) used in tensor predictor regression.

Examples

```
# The dimension of predictor
p <- c(10, 10, 10)
# The envelope dimensions u.
u <- c(1, 1, 1)
# The dimension of response
r <- 5
# The sample size
n <- 200

# Simulate the data with TPRsim.
dat <- TPRsim(p = p, r = r, u = u, n = n)
x <- dat$x
y <- dat$y
B <- dat$coefficients

fit_std <- TPR.fit(x, y, method="standard")
fit_FG <- TPR.fit(x, y, u, method="FG")
fit_pls <- TPR.fit(x, y, u, method="PLS")

rTensor::fnorm(B-stats::coef(fit_std))
rTensor::fnorm(B-stats::coef(fit_FG))
rTensor::fnorm(B-stats::coef(fit_pls))

## ----- Pass a list or an environment to x also works ----- ##
# Pass a list to x
l <- dat[c("x", "y")]
fit_std_l <- TPR.fit(l, method="standard")
```



```

# Pass an environment to x
e <- new.env()
e$x <- dat$x
e$y <- dat$y
fit_std_e <- TPR.fit(e, method="standard")

## ----- Use dataset "square" included in the package ----- ##
data("square")
x <- square$x
y <- square$y
fit_std <- TPR.fit(x, y, method="standard")

```

TPRdim	<i>Envelope dimension by cross-validation for tensor predictor regression (TPR).</i>
--------	--

Description

Select the envelope dimension by cross-validation for tensor predictor regression.

Usage

```
TPRdim(x, y, maxdim = 10, nfolds = 5)
```

Arguments

x	The predictor tensor instance of dimension $p_1 \times p_2 \times \cdots \times p_m \times n$, where n is the sample size. Array with the same dimensions and matrix with dimension $p \times n$ are acceptable.
y	The response matrix of dimension $r \times n$, where n is the sample size. Vector of length n is acceptable.
maxdim	The largest dimension to be considered for selection.
nfolds	Number of folds for cross-validation.

Details

According to Zhang and Li (2017), the dimensions of envelopes at each mode are assumed to be equal, so the u returned is a single value representing the equal envelope dimension.

For each dimension u in $1:\text{maxdim}$, we obtain the prediction

$$\hat{Y}_i = \hat{B}_{(m+1)} \text{vec}(X_i)$$

for each predictor X_i in the k -th testing dataset, $k = 1, \dots, \text{nfolds}$, where \hat{B} is the estimated coefficient based on the k -th training dataset. And the mean squared error for the k -th testing dataset is defined as

$$1/nk \sum_{i=1}^{nk} \|Y_i - \hat{Y}_i\|_F^2,$$

where nk is the sample size of the k -th testing dataset and $\|\cdot\|_F$ denotes the Frobenius norm. Then, the average of n folds mean squared error is recorded as cross-validation mean squared error for the dimension u .

Value

`mincv` The minimal cross-validation mean squared error.
`u` The envelope subspace dimension selected.

References

Zhang, X. and Li, L., 2017. Tensor envelope partial least-squares regression. *Technometrics*, 59(4), pp.426-436.

See Also

[TPRsim](#).

Examples

```
# The dimension of predictor
p <- c(10, 10, 10)
# The envelope dimensions u.
u <- c(1, 1, 1)
# The dimension of response
r <- 5
# The sample size
n <- 200
dat <- TPRsim(p = p, r = r, u = u, n = n)
x <- dat$x
y <- dat$y
TPRdim(x, y, maxdim = 5)

## Use dataset square. (time-consuming)

data("square")
x <- square$x
y <- square$y
# check the dimension of x
dim(x)
# use 32 as the maximal envelope dimension
TPRdim(x, y, maxdim=32)
```

TPRsim *Generate simulation data for tensor predictor regression (TPR)*

Description

This function is used to generate simulation data used in tensor prediction regression.

Usage

TPRsim(p, r, u, n)

Arguments

p	The dimension of predictor, a vector in the form of (p_1, \dots, p_m) .
r	The dimension of response, a scale.
u	The structural dimension of envelopes at each mode, a vector with the same length as p.
n	The sample size.

Details

The tensor predictor regression model is of the form,

$$Y = B_{(m+1)} \text{vec}(X) + \epsilon$$

where response $Y \in R^r$, predictor $X \in R^{p_1 \times \dots \times p_m}$, $B \in R^{p_1 \times \dots \times p_m \times r}$ and the error term is multivariate normal distributed. The predictor is tensor normal distributed,

$$X \sim TN(0; \Sigma_1, \dots, \Sigma_m)$$

According to the tensor envelope structure, we have

$$B = [\Theta; \Gamma_1, \dots, \Gamma_m, I_p],$$

$$\Sigma_k = \Gamma_k \Omega_k \Gamma_k^T + \Gamma_{0k} \Omega_{0k} \Gamma_{0k}^T,$$

for some $\Theta \in R^{u_1 \times \dots \times u_m \times p}$, $\Omega_k \in R^{u_k \times u_k}$ and $\Omega_{0k} \in R^{(p_k - u_k) \times (p_k - u_k)}$, $k = 1, \dots, m$.

Value

x	The predictor of dimension $p_1 \times \dots \times p_m \times n$.
y	The response of dimension $r \times n$.
Gamma	A list of envelope subspace basis of dimension $p_k \times u_k$, $k = 1, \dots, m$.
coefficients	The tensor coefficients of dimension $p_1 \times \dots \times p_m \times r$.
Sigma	A lists of estimated covariance matrices at each mode for the tensor predictors, i.e., $\Sigma_1, \dots, \Sigma_m$.
p, r, u	The input p, r, u.

Note

The length of p must match that of u , and each element of u must be less than the corresponding element in p .

References

Zhang, X. and Li, L., 2017. Tensor envelope partial least-squares regression. *Technometrics*, 59(4), pp.426-436.

See Also

[TPR.fit](#), [TRRsim](#).

Examples

```
p <- c(10, 10, 10)
u <- c(1, 1, 1)
r <- 5
n <- 200
dat <- TPRsim(p = p, r = r, u = u, n = n)
x <- dat$x
y <- dat$y
fit_std <- TPR.fit(x, y, method="standard")
```

TRR.fit

Tensor response regression

Description

This function is used for estimation of tensor response regression. The available method including standard OLS type estimation, PLS type of estimation as well as envelope estimation with FG, 1D and ECD approaches.

Usage

```
TRR.fit(x, y, u, method=c('standard', 'FG', '1D', 'ECD', 'PLS'), Gamma_init = NULL)
```

Arguments

- x The predictor matrix of dimension $p \times n$. Vector of length n is acceptable. If y is missing, x should be a list or an environment consisting of predictor and response datasets.
- y The response tensor instance with dimension $r_1 \times r_2 \times \dots \times r_m \times n$, where n is the sample size. Array with the same dimensions and matrix with dimension $r \times n$ are acceptable.
- u The dimension of envelope subspace. $u = (u_1, \dots, u_m)$. Used for methods "FG", "1D", "ECD" and "PLS". User can use [TRRdim](#) to select dimension.

method	The method used for estimation of tensor response regression. There are four possible choices. <ul style="list-style-type: none"> • "standard": The standard OLS type estimation. • "FG": Envelope estimation with full Grassmannian (FG) algorithm. • "1D": Envelope estimation with one dimensional optimization approaches by 1D algorithm. • "ECD": Envelope estimation with one dimensional optimization approaches by ECD algorithm. • "PLS": The SIMPLS-type estimation without manifold optimization.
Gamma_init	A list specifying the initial envelope subspace basis for "FG" method. By default, the estimators given by "1D" algorithm is used.

Details

Please refer to **Details** part of [TRRsim](#) for the description of the tensor response regression model.

When samples are insufficient, it is possible that the estimation of error covariance matrix Σ is not available. However, if using ordinary least square method (`method = "standard"`), as long as sample covariance matrix of predictor x is nonsingular, `coefficients`, `fitted.values`, `residuals` are still returned.

Value

`TRR.fit` returns an object of class "Tenv".

The function `summary` (i.e., `summary.Tenv`) is used to print the summary of the results, including additional information, e.g., the p-value and the standard error for coefficients, and the prediction mean squared error.

The functions `coefficients`, `fitted.values` and `residuals` can be used to extract different features returned from `TRR.fit`.

The function `plot` (i.e., `plot.Tenv`) plots the two-dimensional coefficients and p-value for object of class "Tenv".

The function `predict` (i.e., `predict.Tenv`) predicts response for the object returned from `TRR.fit` function.

<code>x</code>	The original predictor dataset.
<code>y</code>	The original response dataset.
<code>call</code>	The matched call.
<code>method</code>	The implemented method.
<code>coefficients</code>	The estimation of regression coefficient tensor.
<code>Gamma</code>	The estimation of envelope subspace basis.
<code>Sigma</code>	A lists of estimated covariance matrices at each mode for the error term.
<code>fitted.values</code>	The fitted response tensor.
<code>residuals</code>	The residuals tensor.

References

Li, L. and Zhang, X., 2017. Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.

See Also

[summary.Tenv](#) for summaries, calculating mean squared error from the prediction.

[plot.Tenv](#)(via `graphics::image`) for drawing the two-dimensional coefficient plot and p -value plot.

[predict.Tenv](#) for prediction.

The generic functions [coef](#), [residuals](#), [fitted](#).

[TRRdim](#) for selecting the dimension of envelope by information criteria.

[TRRsim](#) for generating the simulated data used in tensor response regression.

The simulated data [bat](#) used in tensor response regression.

Examples

```
# The dimension of response
r <- c(10, 10, 10)
# The envelope dimensions u.
u <- c(2, 2, 2)
# The dimension of predictor
p <- 5
# The sample size
n <- 100

# Simulate the data with TRRsim.
dat <- TRRsim(r = r, p = p, u = u, n = n)
x <- dat$x
y <- dat$y
B <- dat$coefficients

fit_std <- TRR.fit(x, y, method="standard")
fit_fg <- TRR.fit(x, y, u, method="FG")
fit_1D <- TRR.fit(x, y, u, method="1D")
fit_pls <- TRR.fit(x, y, u, method="PLS")
fit_ECD <- TRR.fit(x, y, u, method="ECD")

rTensor::fnorm(B-stats::coef(fit_std))
rTensor::fnorm(B-stats::coef(fit_fg))
rTensor::fnorm(B-stats::coef(fit_1D))
rTensor::fnorm(B-stats::coef(fit_pls))
rTensor::fnorm(B-stats::coef(fit_ECD))

# Extract the mean squared error, p-value and standard error from summary
summary(fit_std)$mse
summary(fit_std)$p_val
summary(fit_std)$se
```

```
## ----- Pass a list or an environment to x also works ----- ##
# Pass a list to x
l <- dat[c("x", "y")]
fit_std_l <- TRR.fit(l, method="standard")

# Pass an environment to x
e <- new.env()
e$x <- dat$x
e$y <- dat$y
fit_std_e <- TRR.fit(e, method="standard")

## ----- Use dataset "bat" included in the package ----- ##
data("bat")
x <- bat$x
y <- bat$y
fit_std <- TRR.fit(x, y, method="standard")
```

TRRdim

*Envelope dimension selection for tensor response regression (TRR)***Description**

This function uses the 1D-BIC criterion proposed by Zhang, X., & Mai, Q. (2018) to select envelope dimensions in tensor response regression. Refer to [oneD_bic](#) for more details.

Usage

```
TRRdim(x, y, C = NULL, maxdim = 10, ...)
```

Arguments

x	The predictor matrix of dimension $p \times n$. Vector of length n is acceptable.
y	The response tensor instance with dimension $r_1 \times r_2 \times \dots \times r_m \times n$, where n is the sample size. Array with the same dimensions and matrix with dimension $r \times n$ are acceptable.
C	The parameter passed to oneD_bic . Default is $nrow(x) = p$.
maxdim	The maximum envelope dimension to be considered. Default is 10.
...	Additional arguments passed to oneD_bic .

Details

See [oneD_bic](#) for more details on the definition of 1D-BIC criterion and on the arguments C and the additional arguments.

Let B denote the estimated envelope with the selected dimension u , then the prediction is $\hat{Y}_i = B \bar{\times}_{(m+1)} X_i$ for each observation. And the mean squared error is defined as $1/n \sum_{i=1}^n \|Y_i - \hat{Y}_i\|_F^2$, where $\|\cdot\|_F$ denotes the Frobenius norm.

Value

bicval	The minimal BIC values for each mode.
u	The optimal envelope subspace dimension (u_1, u_2, \dots, u_m) .
mse	The prediction mean squared error using the selected envelope basis.

References

- Li, L. and Zhang, X., 2017. Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.
- Zhang, X. and Mai, Q., 2018. Model-free envelope dimension selection. *Electronic Journal of Statistics*, 12(2), pp.2193-2216.

See Also

[oneD_bic](#), [TRRsim](#).

Examples

```
# The dimension of response
r <- c(10, 10, 10)
# The envelope dimensions u.
u <- c(2, 2, 2)
# The dimension of predictor
p <- 5
# The sample size
n <- 100

# Simulate the data with TRRsim.
dat <- TRRsim(r = r, p = p, u = u, n = n)
x <- dat$x
y <- dat$y

TRRdim(x, y) # The estimated envelope dimensions are the same as u.

## Use dataset bat. (time-consuming)

data("bat")
x <- bat$x
y <- bat$y
# check the dimension of y
dim(y)
# use 32 as the maximal envelope dimension
TRRdim(x, y, maxdim=32)
```


TRRsim

*Generate simulation data for tensor response regression (TRR)***Description**

This function is used to generate simulation data used in tensor response regression.

Usage

```
TRRsim(r, p, u, n)
```

Arguments

r	The dimension of response, a vector with length larger than 2.
p	The dimension of predictor, a scale.
u	The structural dimension of envelopes at each mode, a vector with the same length as r.
n	The sample size.

Details

The tensor response regression model is of the form,

$$Y = B \bar{\times}_{(m+1)} X + \epsilon$$

where predictor $X \in R^p$, response $Y \in R^{r_1 \times \dots \times r_m}$, $B \in R^{r_1 \times \dots \times r_m \times p}$ and the error term is tensor normal distributed as follows,

$$\epsilon \sim TN(0; \Sigma_1, \dots, \Sigma_m).$$

According to the tensor envelope structure, we have

$$B = [\Theta; \Gamma_1, \dots, \Gamma_m, I_p],$$

$$\Sigma_k = \Gamma_k \Omega_k \Gamma_k^T + \Gamma_{0k} \Omega_{0k} \Gamma_{0k}^T,$$

for some $\Theta \in R^{u_1 \times \dots \times u_m \times p}$, $\Omega_k \in R^{u_k \times u_k}$ and $\Omega_{0k} \in R^{(p_k - u_k) \times (p_k - u_k)}$, $k = 1, \dots, m$.

Value

x	The predictor of dimension $p \times n$.
y	The response of dimension $r_1 \times \dots \times r_m \times n$.
Gamma	The envelope subspace basis of dimension $r_k \times u_k$, $k = 1, \dots, m$.
coefficients	The tensor coefficients of dimension $r_1 \times \dots \times r_m \times p$.
Sigma	A lists of estimated covariance matrices at each mode for the error term, i.e., $\Sigma_1, \dots, \Sigma_m$.
p, r, u	The input p, r, u.

Note

The length of r must match that of u , and each element of u must be less than the corresponding element in r .

References

Li, L. and Zhang, X., 2017. Parsimonious tensor response regression. *Journal of the American Statistical Association*, 112(519), pp.1131-1146.

See Also

[TPR.fit](#), [TPRsim](#).

Examples

```
r <- c(10, 10, 10)
u <- c(2, 2, 2)
p <- 5
n <- 100
dat <- TRRsim(r = r, p = p, u = u, n = n)
x <- dat$x
y <- dat$y
fit_std <- TRR.fit(x, y, method="standard")
```

 ttt

Matrix product of two tensors

Description

Matrix product of two tensors unfolded on the specified modes.

Usage

```
ttt(x, y, ms)
```

Arguments

x	A tensor instance.
y	A tensor instance.
ms	The indices of the modes to compute on. A single value or a vector.

Details

Suppose x is a s -way tensor with dimension $p_1 \times \dots \times p_s$ and y is a t -way tensor with dimension $r_1 \times \dots \times r_t$. ms specifies the indices on which the tensors x and y are unfolded as columns. Thus, ms must be a subset of $1:\min\{s, t\}$. Meanwhile, the sizes of the dimensions specified by ms must match, e.g., if $ms = 1:k$ where $k \leq \min\{s, t\}$, then $p_1 \times \dots \times p_k = r_1 \times \dots \times r_k$. Let X_0 and Y_0 denote the unfolded matrices, the matrix $X_0 \times Y_0^T$ is returned. See **Examples** for a better illustration.

Value

Return the matrix product of tensors x and y.

Examples

```
x <- rTensor::as.tensor(array(runif(24), c(3, 4, 2)))
y <- rTensor::as.tensor(array(runif(24), c(3, 4, 2)))
z <- ttt(x, y, 1:2)
```

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