

Package: HDTSA (via r-universe)

December 2, 2024

Type Package

Title High Dimensional Time Series Analysis Tools

Version 1.0.5

Date 2024-11-30

Author Jinyuan Chang [aut], Jing He [aut], Chen Lin [aut, cre], Qiwei Yao [aut]

Maintainer Chen Lin <linchen@mail.swufe.edu.cn>

Description An implementation for high-dimensional time series analysis methods, including factor model for vector time series proposed by Lam and Yao (2012) <[doi:10.1214/12-AOS970](https://doi.org/10.1214/12-AOS970)> and Chang, Guo and Yao (2015) <[doi:10.1016/j.jeconom.2015.03.024](https://doi.org/10.1016/j.jeconom.2015.03.024)>, martingale difference test proposed by Chang, Jiang and Shao (2023) <[doi:10.1016/j.jeconom.2022.09.001](https://doi.org/10.1016/j.jeconom.2022.09.001)>, principal component analysis for vector time series proposed by Chang, Guo and Yao (2018) <[doi:10.1214/17-AOS1613](https://doi.org/10.1214/17-AOS1613)>, cointegration analysis proposed by Zhang, Robinson and Yao (2019) <[doi:10.1080/01621459.2018.1458620](https://doi.org/10.1080/01621459.2018.1458620)>, unit root test proposed by Chang, Cheng and Yao (2022) <[doi:10.1093/biomet/asab034](https://doi.org/10.1093/biomet/asab034)>, white noise test proposed by Chang, Yao and Zhou (2017) <[doi:10.1093/biomet/asw066](https://doi.org/10.1093/biomet/asw066)>, CP-decomposition for matrix time series proposed by Chang et al. (2023) <[doi:10.1093/jrsssb/qkac011](https://doi.org/10.1093/jrsssb/qkac011)> and Chang et al. (2024) <[doi:10.48550/arXiv.2410.05634](https://doi.org/10.48550/arXiv.2410.05634)>, and statistical inference for spectral density matrix proposed by Chang et al. (2022) <[doi:10.48550/arXiv.2212.13686](https://doi.org/10.48550/arXiv.2212.13686)>.

License GPL-3

Depends R (>= 3.5.0)

Imports stats, Rcpp, clime, sandwich, methods, MASS, geigen, jointDiag, vars, forecast

LinkingTo RcppEigen, Rcpp

Suggests knitr

NeedsCompilation yes

RoxygenNote 7.3.2

Encoding UTF-8

URL <https://github.com/Linc2021/HDTSA>

BugReports <https://github.com/Linc2021/HDTSA/issues>

Repository CRAN

Date/Publication 2024-12-02 10:00:23 UTC

Config/pak/sysreqs libssl-dev

Contents

Coint	2
CP_MTS	5
DGP.CP	8
Factors	9
FamaFrench	11
HDSReg	12
IPindices	14
MartG_test	15
PCA_TS	17
predict.factors	20
predict.mtsep	21
predict.tspca	23
QWldata	24
SpecMulTest	25
SpecTest	27
UR_test	28
WN_test	29
Index	32

Coint	<i>Identifying the cointegration rank of nonstationary vector time series</i>
-------	---

Description

Coint() deals with cointegration analysis for high-dimensional vector time series proposed in Zhang, Robinson and Yao (2019). Consider the model:

$$\mathbf{y}_t = \mathbf{A}\mathbf{x}_t,$$

where \mathbf{A} is a $p \times p$ unknown and invertible constant matrix, $\mathbf{x}_t = (\mathbf{x}'_{t,1}, \mathbf{x}'_{t,2})'$ is a latent $p \times 1$ process, $\mathbf{x}_{t,2}$ is an $r \times 1$ $I(0)$ process, $\mathbf{x}_{t,1}$ is a process with nonstationary components, and no linear combination of $\mathbf{x}_{t,1}$ is $I(0)$. This function aims to estimate the cointegration rank r and the invertible constant matrix \mathbf{A} .

Usage

```
Coint(
  Y,
  lag.k = 5,
  type = c("acf", "urtest", "both"),
  c0 = 0.3,
  m = 20,
  alpha = 0.01
)
```

Arguments

Y An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$.

lag.k The time lag K used to calculate the nonnegative definite matrix $\hat{\mathbf{W}}_y$:

$$\hat{\mathbf{W}}_y = \sum_{k=0}^K \hat{\Sigma}_y(k) \hat{\Sigma}_y(k)',$$

where $\hat{\Sigma}_y(k)$ is the sample autocovariance of \mathbf{y}_t at lag k . The default is 5.

type The method used to identify the cointegration rank. Available options include: "acf" (the default) for the method based on the sample autocorrelations, "urtest" for the method based on the unit root tests, and "both" to apply these two methods. See Section 2.3 of Zhang, Robinson and Yao (2019) and 'Details' for more information.

c0 The prescribed constant c_0 involved in the method based on the sample correlations, which is used when `type = "acf"` or `type = "both"`. See Section 2.3 of Zhang, Robinson and Yao (2019) and 'Details' for more information. The default is 0.3.

m The prescribed constant m involved in the method based on the sample correlations, which is used when `type = "acf"` or `type = "both"`. See Section 2.3 of Zhang, Robinson and Yao (2019) and 'Details' for more information. The default is 20.

alpha The significance level α of the unit root tests, which is used when `type = "urtest"` or `type = "both"`. See 'Details'. The default is 0.01.

Details

Write $\hat{\mathbf{x}}_t = \hat{\mathbf{A}}' \mathbf{y}_t \equiv (\hat{x}_t^1, \dots, \hat{x}_t^p)'$. When `type = "acf"`, `Coint()` estimates r by

$$\hat{r} = \sum_{i=1}^p 1 \left\{ \frac{S_i(m)}{m} < c_0 \right\}$$

for some constant $c_0 \in (0, 1)$ and some large constant m , where $S_i(m)$ is the sum of the sample autocorrelations of \hat{x}_t^i over lags 1 to m , which is specified in Section 2.3 of Zhang, Robinson and Yao (2019).

When type = "urtest", Coint() estimates r by unit root tests. For $i = 1, \dots, p$, consider the null hypothesis

$$H_{0,i} : \hat{x}_t^{p-i+1} \sim I(0).$$

The estimation procedure for r can be implemented as follows:

Step 1. Start with $i = 1$. Perform the unit root test proposed in Chang, Cheng and Yao (2021) for $H_{0,i}$.

Step 2. If the null hypothesis is not rejected at the significance level α , increment i by 1 and repeat Step 1. Otherwise, stop the procedure and denote the value of i at termination as i_0 . The cointegration rank is then estimated as $\hat{r} = i_0 - 1$.

Value

An object of class "coint", which contains the following components:

A	The estimated $\hat{\mathbf{A}}$.
coint_rank	The estimated cointegration rank \hat{r} .
lag.k	The time lag used in function.
method	A string indicating which method is used to identify the cointegration rank.

References

- Chang, J., Cheng, G., & Yao, Q. (2022). Testing for unit roots based on sample autocovariances. *Biometrika*, **109**, 543–550. doi:10.1093/biomet/asab034.
- Zhang, R., Robinson, P., & Yao, Q. (2019). Identifying cointegration by eigenanalysis. *Journal of the American Statistical Association*, **114**, 916–927. doi:10.1080/01621459.2018.1458620.

Examples

```
# Example 1 (Example 1 in Zhang, Robinson and Yao (2019))
## Generate yt
p <- 10
n <- 1000
r <- 3
d <- 1
X <- mat.or.vec(p, n)
X[1,] <- arima.sim(n-d, model = list(order=c(0, d, 0)))
for(i in 2:3)X[i,] <- rnorm(n)
for(i in 4:(r+1)) X[i, ] <- arima.sim(model = list(ar = 0.5), n)
for(i in (r+2):p) X[i, ] <- arima.sim(n = (n-d), model = list(order=c(1, d, 1), ar=0.6, ma=0.8))
M1 <- matrix(c(1, 1, 0, 1/2, 0, 1, 0, 1, 0), ncol = 3, byrow = TRUE)
A <- matrix(runif(p*p, -3, 3), ncol = p)
A[1:3,1:3] <- M1
Y <- t(A%%X)

Coint(Y, type = "both")
```

Description

CP_MTS() deals with the estimation of the CP-factor model for matrix time series:

$$\mathbf{Y}_t = \mathbf{A}\mathbf{X}_t\mathbf{B}' + \boldsymbol{\epsilon}_t,$$

where $\mathbf{X}_t = \text{diag}(x_{t,1}, \dots, x_{t,d})$ is a $d \times d$ unobservable diagonal matrix, $\boldsymbol{\epsilon}_t$ is a $p \times q$ matrix white noise, \mathbf{A} and \mathbf{B} are, respectively, $p \times d$ and $q \times d$ unknown constant matrices with their columns being unit vectors, and $1 \leq d < \min(p, q)$ is an unknown integer. Let $\text{rank}(\mathbf{A}) = d_1$ and $\text{rank}(\mathbf{B}) = d_2$ with some unknown $d_1, d_2 \leq d$. This function aims to estimate d, d_1, d_2 and the loading matrices \mathbf{A} and \mathbf{B} using the methods proposed in Chang et al. (2023) and Chang et al. (2024).

Usage

```
CP_MTS(
  Y,
  xi = NULL,
  Rank = NULL,
  lag.k = 20,
  lag.ktilde = 10,
  method = c("CP.Direct", "CP.Refined", "CP.Unified"),
  thresh1 = FALSE,
  thresh2 = FALSE,
  thresh3 = FALSE,
  delta1 = 2 * sqrt(log(dim(Y)[2] * dim(Y)[3])/dim(Y)[1]),
  delta2 = delta1,
  delta3 = delta1
)
```

Arguments

- | | |
|------|---|
| Y | An $n \times p \times q$ array, where n is the number of observations of the $p \times q$ matrix time series $\{\mathbf{Y}_t\}_{t=1}^n$. |
| xi | An $n \times 1$ vector $\boldsymbol{\xi} = (\xi_1, \dots, \xi_n)'$, where ξ_t represents a linear combination of \mathbf{Y}_t . If xi = NULL (the default), ξ_t is determined by the PCA method introduced in Section 5.1 of Chang et al. (2023). Otherwise, xi can be given by the users. |
| Rank | A list containing the following components: d representing the number of columns of \mathbf{A} and \mathbf{B} , d1 representing the rank of \mathbf{A} , and d2 representing the rank of \mathbf{B} . If set to NULL (default), $d, d_1,$ and d_2 will be estimated. Otherwise, they can be given by the users. |

lag.k The time lag K used to calculate the nonnegative definite matrices $\hat{\mathbf{M}}_1$ and $\hat{\mathbf{M}}_2$ when method = "CP.Refined" or method = "CP.Unified":

$$\hat{\mathbf{M}}_1 = \sum_{k=1}^K \hat{\Sigma}_k \hat{\Sigma}_k' \text{ and } \hat{\mathbf{M}}_2 = \sum_{k=1}^K \hat{\Sigma}_k' \hat{\Sigma}_k,$$

where $\hat{\Sigma}_k$ is an estimate of the cross-covariance between \mathbf{Y}_t and ξ_t at lag k . See 'Details'. The default is 20.

lag.ktilde The time lag \tilde{K} involved in the unified estimation method [See (16) in Chang et al. (2024)], which is used when method = "CP.Unified". The default is 10.

method A string indicating which CP-decomposition method is used. Available options include: "CP.Direct" (the default) for the direct estimation method [See Section 3.1 of Chang et al. (2023)], "CP.Refined" for the refined estimation method [See Section 3.2 of Chang et al. (2023)], and "CP.Unified" for the unified estimation method [See Section 4 of Chang et al. (2024)]. The validity of methods "CP.Direct" and "CP.Refined" depends on the assumption $d_1 = d_2 = d$. When $d_1, d_2 \leq d$, the method "CP.Unified" can be applied. See Chang et al. (2024) for details.

thresh1 Logical. If FALSE (the default), no thresholding will be applied in $\hat{\Sigma}_k$, which indicates that the threshold level $\delta_1 = 0$. If TRUE, δ_1 will be set through delta1. thresh1 is used for all three methods. See 'Details'.

thresh2 Logical. If FALSE (the default), no thresholding will be applied in $\check{\Sigma}_k$, which indicates that the threshold level $\delta_2 = 0$. If TRUE, δ_2 will be set through delta2. thresh2 is used only when method = "CP.Refined". See 'Details'.

thresh3 Logical. If FALSE (the default), no thresholding will be applied in $\vec{\Sigma}_k$, which indicates that the threshold level $\delta_3 = 0$. If TRUE, δ_3 will be set through delta3. thresh3 is used only when method = "CP.Unified". See 'Details'.

delta1 The value of the threshold level δ_1 . The default is $\delta_1 = 2\sqrt{n^{-1} \log(pq)}$.

delta2 The value of the threshold level δ_2 . The default is $\delta_2 = 2\sqrt{n^{-1} \log(pq)}$.

delta3 The value of the threshold level δ_3 . The default is $\delta_3 = 2\sqrt{n^{-1} \log(pq)}$.

Details

All three CP-decomposition methods involve the estimation of the autocovariance of \mathbf{Y}_t and ξ_t at lag k , which is defined as follows:

$$\hat{\Sigma}_k = T_{\delta_1} \{ \hat{\Sigma}_{\mathbf{Y}, \xi}(k) \} \text{ with } \hat{\Sigma}_{\mathbf{Y}, \xi}(k) = \frac{1}{n-k} \sum_{t=k+1}^n (\mathbf{Y}_t - \bar{\mathbf{Y}})(\xi_{t-k} - \bar{\xi}),$$

where $\bar{\mathbf{Y}} = n^{-1} \sum_{t=1}^n \mathbf{Y}_t$, $\bar{\xi} = n^{-1} \sum_{t=1}^n \xi_t$ and $T_{\delta_1}(\cdot)$ is a threshold operator defined as $T_{\delta_1}(\mathbf{W}) = \{w_{i,j} 1(|w_{i,j}| \geq \delta_1)\}$ for any matrix $\mathbf{W} = (w_{i,j})$, with the threshold level $\delta_1 \geq 0$ and $1(\cdot)$ representing the indicator function. Chang et al. (2023) and Chang et al. (2024) suggest to choose $\delta_1 = 0$ when p, q are fixed and $\delta_1 > 0$ when $pq \gg n$.

The refined estimation method involves

$$\check{\Sigma}_k = T_{\delta_2} \{ \hat{\Sigma}_{\bar{\mathbf{Y}}}(k) \} \text{ with } \hat{\Sigma}_{\bar{\mathbf{Y}}}(k) = \frac{1}{n-k} \sum_{t=k+1}^n (\mathbf{Y}_t - \bar{\mathbf{Y}}) \otimes \text{vec}(\mathbf{Y}_{t-k} - \bar{\mathbf{Y}}),$$

where $T_{\delta_2}(\cdot)$ is a threshold operator with the threshold level $\delta_2 \geq 0$, and $\text{vec}(\cdot)$ is a vectorization operator with $\text{vec}(\mathbf{H})$ being the $(m_1 m_2) \times 1$ vector obtained by stacking the columns of the $m_1 \times m_2$ matrix \mathbf{H} . See Section 3.2.2 of Chang et al. (2023) for details.

The unified estimation method involves

$$\vec{\Sigma}_k = T_{\delta_3}\{\hat{\Sigma}_{\bar{\mathbf{Y}}}(k)\} \text{ with } \hat{\Sigma}_{\bar{\mathbf{Y}}}(k) = \frac{1}{n-k} \sum_{t=k+1}^n \text{vec}(\mathbf{Y}_t - \bar{\mathbf{Y}})\{\text{vec}(\mathbf{Y}_{t-k} - \bar{\mathbf{Y}})\}',$$

where $T_{\delta_3}(\cdot)$ is a threshold operator with the threshold level $\delta_3 \geq 0$. See Section 4.2 of Chang et al. (2024) for details.

Value

An object of class "mtscp", which contains the following components:

A	The estimated $p \times \hat{d}$ left loading matrix $\hat{\mathbf{A}}$.
B	The estimated $q \times \hat{d}$ right loading matrix $\hat{\mathbf{B}}$.
f	The estimated latent process $\hat{x}_{t,1}, \dots, \hat{x}_{t,\hat{d}}$.
Rank	The estimated \hat{d}_1, \hat{d}_2 , and \hat{d} .
method	A string indicating which CP-decomposition method is used.

References

- Chang, J., Du, Y., Huang, G., & Yao, Q. (2024). Identification and estimation for matrix time series CP-factor models. *arXiv preprint*. doi:10.48550/arXiv.2410.05634.
- Chang, J., He, J., Yang, L., & Yao, Q. (2023). Modelling matrix time series via a tensor CP-decomposition. *Journal of the Royal Statistical Society Series B: Statistical Methodology*, **85**, 127–148. doi:10.1093/jrsssb/qqac011.

Examples

```
# Example 1.
p <- 10
q <- 10
n <- 400
d = d1 = d2 <- 3
## DGP.CP() generates simulated data for the example in Chang et al. (2024).
data <- DGP.CP(n, p, q, d, d1, d2)
Y <- data$Y

## d is unknown
res1 <- CP_MTS(Y, method = "CP.Direct")
res2 <- CP_MTS(Y, method = "CP.Refined")
res3 <- CP_MTS(Y, method = "CP.Unified")

## d is known
res4 <- CP_MTS(Y, Rank = list(d = 3), method = "CP.Direct")
res5 <- CP_MTS(Y, Rank = list(d = 3), method = "CP.Refined")
```

```

# Example 2.
p <- 10
q <- 10
n <- 400
d1 = d2 <- 2
d <-3
data <- DGP.CP(n, p, q, d, d1, d2)
Y1 <- data$Y

## d, d1 and d2 are unknown
res6 <- CP_MTS(Y1, method = "CP.Unified")
## d, d1 and d2 are known
res7 <- CP_MTS(Y1, Rank = list(d = 3, d1 = 2, d2 = 2), method = "CP.Unified")

```

DGP.CP

Generating simulated data for the example in Chang et al. (2024)

Description

DGP.CP() function generates simulated data following the data generating process described in Section 7.1 of Chang et al. (2024).

Usage

```
DGP.CP(n, p, q, d, d1, d2)
```

Arguments

n	Integer. The number of observations of the $p \times q$ matrix time series \mathbf{Y}_t .
p	Integer. The number of rows of \mathbf{Y}_t .
q	Integer. The number of columns of \mathbf{Y}_t .
d	Integer. The number of columns of the factor loading matrices \mathbf{A} and \mathbf{B} .
d1	Integer. The rank of the $p \times d$ matrix \mathbf{A} .
d2	Integer. The rank of the $q \times d$ matrix \mathbf{B} .

Details

We generate

$$\mathbf{Y}_t = \mathbf{A}\mathbf{X}_t\mathbf{B}' + \boldsymbol{\epsilon}_t$$

for any $t = 1, \dots, n$, where $\mathbf{X}_t = \text{diag}(\mathbf{x}_t)$ with $\mathbf{x}_t = (x_{t,1}, \dots, x_{t,d})'$ being a $d \times 1$ time series, $\boldsymbol{\epsilon}_t$ is a $p \times q$ matrix white noise, and \mathbf{A} and \mathbf{B} are, respectively, $p \times d$ and $q \times d$ factor loading matrices. \mathbf{A} , \mathbf{X}_t , and \mathbf{B} are generated based on the data generating process described in Section 7.1 of Chang et al. (2024) and satisfy $\text{rank}(\mathbf{A}) = d_1$ and $\text{rank}(\mathbf{B}) = d_2$, $1 \leq d_1, d_2 \leq d$.

Value

A list containing the following components:

Y	An $n \times p \times q$ array.
A	The $p \times d$ left loading matrix A .
B	The $q \times d$ right loading matrix B .
X	An $n \times d \times d$ array.

References

Chang, J., Du, Y., Huang, G., & Yao, Q. (2024). Identification and estimation for matrix time series CP-factor models. *arXiv preprint*. doi:10.48550/arXiv.2410.05634.

See Also

[CP_MTS](#).

Examples

```
p <- 10
q <- 10
n <- 400
d = d1 = d2 <- 3
data <- DGP.CP(n,p,q,d1,d2,d)
Y <- data$Y

## The first observation: Y_1
Y[1, , ]
```

Description

Factors() deals with factor modeling for high-dimensional time series proposed in Lam and Yao (2012):

$$\mathbf{y}_t = \mathbf{A}\mathbf{x}_t + \boldsymbol{\epsilon}_t,$$

where \mathbf{x}_t is an $r \times 1$ latent process with (unknown) $r \leq p$, **A** is a $p \times r$ unknown constant matrix, and $\boldsymbol{\epsilon}_t$ is a vector white noise process. The number of factors r and the factor loadings **A** can be estimated in terms of an eigenanalysis for a nonnegative definite matrix, and is therefore applicable when the dimension of \mathbf{y}_t is on the order of a few thousands. This function aims to estimate the number of factors r and the factor loading matrix **A**.

Usage

```
Factors(
  Y,
  lag.k = 5,
  thresh = FALSE,
  delta = 2 * sqrt(log(ncol(Y))/nrow(Y)),
  twostep = FALSE
)
```

Arguments

Y An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$.

lag.k The time lag K used to calculate the nonnegative definite matrix $\hat{\mathbf{M}}$:

$$\hat{\mathbf{M}} = \sum_{k=1}^K T_\delta\{\hat{\Sigma}_y(k)\}T_\delta\{\hat{\Sigma}_y(k)\}',$$

where $\hat{\Sigma}_y(k)$ is the sample autocovariance of \mathbf{y}_t at lag k and $T_\delta(\cdot)$ is a threshold operator with the threshold level $\delta \geq 0$. See 'Details'. The default is 5.

thresh Logical. If **thresh** = FALSE (the default), no thresholding will be applied to estimate $\hat{\mathbf{M}}$. If **thresh** = TRUE, δ will be set through **delta**.

delta The value of the threshold level δ . The default is $\delta = 2\sqrt{n^{-1} \log p}$.

twostep Logical. If **twostep** = FALSE (the default), the standard procedure [See Section 2.2 in Lam and Yao (2012)] for estimating r and \mathbf{A} will be implemented. If **twostep** = TRUE, the two-step estimation procedure [See Section 4 in Lam and Yao (2012)] for estimating r and \mathbf{A} will be implemented.

Details

The threshold operator $T_\delta(\cdot)$ is defined as $T_\delta(\mathbf{W}) = \{w_{i,j}1(|w_{i,j}| \geq \delta)\}$ for any matrix $\mathbf{W} = (w_{i,j})$, with the threshold level $\delta \geq 0$ and $1(\cdot)$ representing the indicator function. We recommend to choose $\delta = 0$ when p is fixed and $\delta > 0$ when $p \gg n$.

Value

An object of class "factors", which contains the following components:

factor_num The estimated number of factors \hat{r} .

loading.mat The estimated $p \times \hat{r}$ factor loading matrix $\hat{\mathbf{A}}$.

X The $n \times \hat{r}$ matrix $\hat{\mathbf{X}} = (\hat{\mathbf{x}}_1, \dots, \hat{\mathbf{x}}_n)'$ with $\hat{\mathbf{x}}_t = \hat{\mathbf{A}}'\hat{\mathbf{y}}_t$.

lag.k The time lag used in function.

References

Lam, C., & Yao, Q. (2012). Factor modelling for high-dimensional time series: Inference for the number of factors. *The Annals of Statistics*, **40**, 694–726. doi:10.1214/12AOS970.

Examples

```
# Example 1 (Example in Section 3.3 of lam and Yao 2012)
## Generate y_t
p <- 200
n <- 400
r <- 3
X <- mat.or.vec(n, r)
A <- matrix(runif(p*r, -1, 1), ncol=r)
x1 <- arima.sim(model=list(ar=c(0.6)), n=n)
x2 <- arima.sim(model=list(ar=c(-0.5)), n=n)
x3 <- arima.sim(model=list(ar=c(0.3)), n=n)
eps <- matrix(rnorm(n*p), p, n)
X <- t(cbind(x1, x2, x3))
Y <- A %*% X + eps
Y <- t(Y)

fac <- Factors(Y,lag.k=2)
r_hat <- fac$factor_num
loading_Mat <- fac$loading.mat
```

FamaFrench

*Fama-French 10*10 return series*

Description

The portfolios are constructed by the intersections of 10 levels of size, denoted by S_1, \dots, S_{10} , and 10 levels of the book equity to market equity ratio (BE), denoted by BE_1, \dots, BE_{10} . The dataset consists of monthly returns from January 1964 to December 2021, which contains 69600 observations for 696 total months.

Usage

```
data(FamaFrench)
```

Format

A data frame with 696 rows and 102 columns. The first column represents the month, and the second column named MKT.RF represents the monthly market returns. The rest of the columns represent the return series for different sizes and BE-ratios.

Source

http://mba.tuck.dartmouth.edu/pages/faculty/ken.french/data_library.html

Description

HDSReg() considers a multivariate time series model which represents a high-dimensional vector process as a sum of three terms: a linear regression of some observed regressors, a linear combination of some latent and serially correlated factors, and a vector white noise:

$$\mathbf{y}_t = \mathbf{D}\mathbf{z}_t + \mathbf{A}\mathbf{x}_t + \boldsymbol{\epsilon}_t,$$

where \mathbf{y}_t and \mathbf{z}_t are, respectively, observable $p \times 1$ and $m \times 1$ time series, \mathbf{x}_t is an $r \times 1$ latent factor process, $\boldsymbol{\epsilon}_t$ is a vector white noise process, \mathbf{D} is an unknown regression coefficient matrix, and \mathbf{A} is an unknown factor loading matrix. This procedure proposed in Chang, Guo and Yao (2015) aims to estimate the regression coefficient matrix \mathbf{D} , the number of factors r and the factor loading matrix \mathbf{A} .

Usage

```
HDSReg(
  Y,
  Z,
  D = NULL,
  lag.k = 5,
  thresh = FALSE,
  delta = 2 * sqrt(log(ncol(Y))/nrow(Y)),
  twostep = FALSE
)
```

Arguments

- | | |
|-------|---|
| Y | An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$. |
| Z | An $n \times m$ data matrix $\mathbf{Z} = (\mathbf{z}_1, \dots, \mathbf{z}_n)'$ consisting of the observed regressors. |
| D | A $p \times m$ regression coefficient matrix $\tilde{\mathbf{D}}$. If D = NULL (the default), our procedure will estimate \mathbf{D} first and let $\tilde{\mathbf{D}}$ be the estimate of \mathbf{D} . If D is given by the users, then $\tilde{\mathbf{D}} = \mathbf{D}$. |
| lag.k | The time lag K used to calculate the nonnegative definite matrix $\hat{\mathbf{M}}_\eta$: |

$$\hat{\mathbf{M}}_\eta = \sum_{k=1}^K T_\delta\{\hat{\boldsymbol{\Sigma}}_\eta(k)\}T_\delta\{\hat{\boldsymbol{\Sigma}}_\eta(k)\}',$$

where $\hat{\boldsymbol{\Sigma}}_\eta(k)$ is the sample autocovariance of $\boldsymbol{\eta}_t = \mathbf{y}_t - \tilde{\mathbf{D}}\mathbf{z}_t$ at lag k and $T_\delta(\cdot)$ is a threshold operator with the threshold level $\delta \geq 0$. See 'Details'. The default is 5.

thresh	Logical. If thresh = FALSE (the default), no thresholding will be applied to estimate $\hat{\mathbf{M}}_\eta$. If thresh = TRUE, δ will be set through delta. See 'Details'.
delta	The value of the threshold level δ . The default is $\delta = 2\sqrt{n^{-1} \log p}$.
twostep	Logical. The same as the argument twostep in Factors .

Details

The threshold operator $T_\delta(\cdot)$ is defined as $T_\delta(\mathbf{W}) = \{w_{i,j}1(|w_{i,j}| \geq \delta)\}$ for any matrix $\mathbf{W} = (w_{i,j})$, with the threshold level $\delta \geq 0$ and $1(\cdot)$ representing the indicator function. We recommend to choose $\delta = 0$ when p is fixed and $\delta > 0$ when $p \gg n$.

Value

An object of class "factors", which contains the following components:

factor_num	The estimated number of factors \hat{r} .
reg.coff.mat	The estimated $p \times m$ regression coefficient matrix $\tilde{\mathbf{D}}$.
loading.mat	The estimated $p \times \hat{r}$ factor loading matrix $\hat{\mathbf{A}}$.
X	The $n \times \hat{r}$ matrix $\tilde{\mathbf{X}} = (\hat{\mathbf{x}}_1, \dots, \hat{\mathbf{x}}_n)'$ with $\hat{\mathbf{x}}_t = \hat{\mathbf{A}}'(y_t - \tilde{\mathbf{D}}z_t)$.
lag.k	The time lag used in function.

References

Chang, J., Guo, B., & Yao, Q. (2015). High dimensional stochastic regression with latent factors, endogeneity and nonlinearity. *Journal of Econometrics*, **189**, 297–312. doi:10.1016/j.jeconom.2015.03.024.

See Also

[Factors](#).

Examples

```
# Example 1 (Example 1 in Chang, Guo and Yao (2015)).
## Generate xt
n <- 400
p <- 200
m <- 2
r <- 3
X <- mat.or.vec(n,r)
x1 <- arima.sim(model = list(ar = c(0.6)), n = n)
x2 <- arima.sim(model = list(ar = c(-0.5)), n = n)
x3 <- arima.sim(model = list(ar = c(0.3)), n = n)
X <- cbind(x1, x2, x3)
X <- t(X)

## Generate yt
Z <- mat.or.vec(m,n)
S1 <- matrix(c(5/8, 1/8, 1/8, 5/8), 2, 2)
Z[,1] <- c(rnorm(m))
```

```

for(i in c(2:n)){
  Z[,i] <- S1%*%Z[, i-1] + c(rnorm(m))
}
D <- matrix(runif(p*m, -2, 2), ncol = m)
A <- matrix(runif(p*r, -2, 2), ncol = r)
eps <- mat.or.vec(n, p)
eps <- matrix(rnorm(n*p), p, n)
Y <- D %*% Z + A %*% X + eps
Y <- t(Y)
Z <- t(Z)

## D is known
res1 <- HDSReg(Y, Z, D, lag.k = 2)
## D is unknown
res2 <- HDSReg(Y, Z, lag.k = 2)

```

IPindices

U.S. Industrial Production indices

Description

The dataset consists of 7 monthly U.S. Industrial Production indices, namely *the total index, nonindustrial supplies, final products, manufacturing, materials, mining, and utilities*, from January 1947 to December 2023 published by the U.S. Federal Reserve.

Usage

```
data(IPindices)
```

Format

A data frame with 924 rows and 8 variables:

DATE The observation date

INDPRO The total index

IPB54000S Nonindustrial supplies

IPFINAL Final products

IPMANSICS Manufacturing

IPMAT Materials

IPMINE Mining

IPUTIL Utilities

Source

<https://fred.stlouisfed.org/release/tables?rid=13&eid=49670>

 MartG_test

Testing for martingale difference hypothesis in high dimension

Description

MartG_test() implements a new test proposed in Chang, Jiang and Shao (2023) for the following hypothesis testing problem:

$$H_0 : \{\mathbf{y}_t\}_{t=1}^n \text{ is a MDS versus } H_1 : \{\mathbf{y}_t\}_{t=1}^n \text{ is not a MDS,}$$

where MDS is the abbreviation of "martingale difference sequence".

Usage

```
MartG_test(
  Y,
  lag.k = 2,
  B = 1000,
  type = c("Linear", "Quad"),
  alpha = 0.05,
  kernel.type = c("QS", "Par", "Bart")
)
```

Arguments

Y	An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$.
lag.k	The time lag K used to calculate the test statistic [See (3) in Chang, Jiang and Shao (2023)]. The default is 2.
B	The number of bootstrap replications for generating multivariate normally distributed random vectors when calculating the critical value. The default is 1000.
type	The map used for constructing the test statistic. Available options include: "Linear" (the default) for the linear identity map and "Quad" for the map including both linear and quadratic terms. type can also be set by the users. See 'Details' and Section 2.1 of Chang, Jiang and Shao (2023) for more information.
alpha	The significance level of the test. The default is 0.05.
kernel.type	The option for choosing the symmetric kernel used in the estimation of long-run covariance matrix. Available options include: "QS" (the default) for the Quadratic spectral kernel, "Par" for the Parzen kernel, and "Bart" for the Bartlett kernel. See Chang, Jiang and Shao (2023) for more information.

Details

Write $\mathbf{x} = (x_1, \dots, x_p)'$. When type = "Linear", the linear identity map is defined as $\phi(\mathbf{x}) = \mathbf{x}$.

When type = "Quad", $\phi(\mathbf{x}) = \{\mathbf{x}', (\mathbf{x}^2)'\}'$ includes both linear and quadratic terms, where $\mathbf{x}^2 = (x_1^2, \dots, x_p^2)'$.

We can also choose $\phi(\mathbf{x}) = \cos(\mathbf{x})$ to capture certain type of nonlinear dependence, where $\cos(\mathbf{x}) = (\cos x_1, \dots, \cos x_p)'$.

See 'Examples'.

Value

An object of class "hdtstest", which contains the following components:

statistic	The test statistic of the test.
p.value	The p-value of the test.
lag.k	The time lag used in function.
type	The map used in function.
kernel.type	The kernel used in function.

References

Chang, J., Jiang, Q., & Shao, X. (2023). Testing the martingale difference hypothesis in high dimension. *Journal of Econometrics*, **235**, 972–1000. doi:10.1016/j.jeconom.2022.09.001.

Examples

```
# Example 1
n <- 200
p <- 10
X <- matrix(rnorm(n*p),n,p)

res <- MartG_test(X, type="Linear")
res <- MartG_test(X, type=cbind(X, X^2)) #the same as type = "Quad"

## map can also be defined as an expression in R.
res <- MartG_test(X, type=quote(cbind(X, X^2))) # expr using quote()
res <- MartG_test(X, type=substitute(cbind(X, X^2))) # expr using substitute()
res <- MartG_test(X, type=expression(cbind(X, X^2))) # expr using expression()
res <- MartG_test(X, type=parse(text="cbind(X, X^2)")) # expr using parse()

## map can also be defined as a function in R.
map_fun <- function(X) {X <- cbind(X, X^2); X}

res <- MartG_test(X, type=map_fun)
Pvalue <- res$p.value
rej <- res$reject
```


Description

PCA_TS() seeks for a contemporaneous linear transformation for a multivariate time series such that the transformed series is segmented into several lower-dimensional subseries:

$$\mathbf{y}_t = \mathbf{A}\mathbf{x}_t,$$

where \mathbf{x}_t is an unobservable $p \times 1$ weakly stationary time series consisting of $q (\geq 1)$ both contemporaneously and serially uncorrelated subseries. See Chang, Guo and Yao (2018).

Usage

```
PCA_TS(  
  Y,  
  lag.k = 5,  
  opt = 1,  
  permutation = c("max", "fdr"),  
  thresh = FALSE,  
  delta = 2 * sqrt(log(ncol(Y))/nrow(Y)),  
  prewhiten = TRUE,  
  m = NULL,  
  beta,  
  control = list()  
)
```

Arguments

Y An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$. The procedure will first normalize \mathbf{y}_t as $\hat{\mathbf{V}}^{-1/2}\mathbf{y}_t$, where $\hat{\mathbf{V}}$ is an estimator for covariance of \mathbf{y}_t . See details below for the selection of $\hat{\mathbf{V}}^{-1}$.

lag.k The time lag K used to calculate the nonnegative definite matrix $\hat{\mathbf{W}}_y$:

$$\hat{\mathbf{W}}_y = \mathbf{I}_p + \sum_{k=1}^K T_\delta\{\hat{\Sigma}_y(k)\}T_\delta\{\hat{\Sigma}_y(k)\}',$$

where $\hat{\Sigma}_y(k)$ is the sample autocovariance of $\hat{\mathbf{V}}^{-1/2}\mathbf{y}_t$ at lag k and $T_\delta(\cdot)$ is a threshold operator with the threshold level $\delta \geq 0$. See 'Details'. The default is 5.

opt An option used to choose which method will be implemented to get a consistent estimate $\hat{\mathbf{V}}$ (or $\hat{\mathbf{V}}^{-1}$) for the covariance (precision) matrix of \mathbf{y}_t . If $\text{opt} = 1$, $\hat{\mathbf{V}}$ will be defined as the sample covariance matrix. If $\text{opt} = 2$, the precision matrix $\hat{\mathbf{V}}^{-1}$ will be calculated by using the function `clime()` of **clime** (Cai, Liu and Luo, 2011) with the arguments passed by `control`.

permutation	The method of permutation procedure to assign the components of $\hat{\mathbf{z}}_t$ to different groups [See Section 2.2.1 in Chang, Guo and Yao (2018)]. Available options include: "max" (the default) for the maximum cross correlation method and "fdr" for the false discovery rate procedure based on multiple tests. See Sections 2.2.2 and 2.2.3 in Chang, Guo and Yao (2018) for more information.
thresh	Logical. If thresh = FALSE (the default), no thresholding will be applied to estimate $\hat{\mathbf{W}}_y$. If thresh = TRUE, the argument delta is used to specify the threshold level δ .
delta	The value of the threshold level δ . The default is $\delta = 2\sqrt{n^{-1} \log p}$.
prewhiten	Logical. If TRUE (the default), we prewhiten each transformed component series of $\hat{\mathbf{z}}_t$ [See Section 2.2.1 in Chang, Guo and Yao (2018)] by fitting a univariate AR model with the order between 0 and 5 determined by AIC. If FALSE, then the prewhiten procedure will not be performed.
m	A positive integer used in the permutation procedure [See (2.10) in Chang, Guo and Yao (2018)]. The default is 10.
beta	The error rate used in the permutation procedure [See (2.16) in Chang, Guo and Yao (2018)] when permutation = "fdr".
control	A list of control arguments. See 'Details'.

Details

The threshold operator $T_\delta(\cdot)$ is defined as $T_\delta(\mathbf{W}) = \{w_{i,j}1(|w_{i,j}| \geq \delta)\}$ for any matrix $\mathbf{W} = (w_{i,j})$, with the threshold level $\delta \geq 0$ and $1(\cdot)$ representing the indicator function. We recommend to choose $\delta = 0$ when p is fixed and $\delta > 0$ when $p \gg n$.

For large p , since the sample covariance matrix may not be consistent, we recommend to use the method proposed in Cai, Liu and Luo (2011) to estimate the precision matrix $\hat{\mathbf{V}}^{-1}$ (opt = 2).

control is a list of arguments passed to the function `clime()`, which contains the following components:

- `nlambda`: Number of values for program generated lambda. The default is 100.
- `lambda.max`: Maximum value of program generated lambda. The default is 0.8.
- `lambda.min`: Minimum value of program generated lambda. The default is 10^{-4} ($n > p$) or 10^{-2} ($n < p$).
- `standardize`: Logical. If `standardize = TRUE`, the variables will be standardized to have mean zero and unit standard deviation. The default is FALSE.
- `linsolver`: An option used to choose which method should be employed. Available options include "primaldual" (the default) and "simplex". Rule of thumb: "primaldual" for large p , "simplex" for small p .

Value

An object of class "tspca", which contains the following components:

B	The $p \times p$ transformation matrix $\hat{\mathbf{B}} = \hat{\mathbf{\Gamma}}_y' \hat{\mathbf{V}}^{-1/2}$, where $\hat{\mathbf{\Gamma}}_y$ is a $p \times p$ orthogonal matrix with the columns being the eigenvectors of $\hat{\mathbf{W}}_y$.
---	--

X	The $n \times p$ matrix $\hat{\mathbf{X}} = (\hat{\mathbf{x}}_1, \dots, \hat{\mathbf{x}}_n)'$ with $\hat{\mathbf{x}}_t = \hat{\mathbf{B}}\mathbf{y}_t$.
NoGroups	The number of groups.
No_of_Members	The number of members in each group.
Groups	The indices of the components of $\hat{\mathbf{x}}_t$ that belong to each group.
method	A string indicating which permutation procedure is performed.

References

- Cai, T., Liu, W., & Luo, X. (2011). A constrained L1 minimization approach for sparse precision matrix estimation. *Journal of the American Statistical Association*, **106**, 594–607. doi:10.1198/jasa.2011.tm10155.
- Chang, J., Guo, B., & Yao, Q. (2018). Principal component analysis for second-order stationary vector time series. *The Annals of Statistics*, **46**, 2094–2124. doi:10.1214/17AOS1613.

Examples

```
# Example 1 (Example 1 in the supplementary material of Chang, Guo and Yao (2018)).
# p=6, x_t consists of 3 independent subseries with 3, 2 and 1 components.

## Generate x_t
p <- 6;n <- 1500
X <- mat.or.vec(p,n)
x <- arima.sim(model = list(ar = c(0.5, 0.3), ma = c(-0.9, 0.3, 1.2,1.3)),
n = n+2, sd = 1)
for(i in 1:3) X[i,] <- x[i:(n+i-1)]
x <- arima.sim(model = list(ar = c(0.8,-0.5),ma = c(1,0.8,1.8) ), n = n+1, sd = 1)
for(i in 4:5) X[i,] <- x[(i-3):(n+i-4)]
x <- arima.sim(model = list(ar = c(-0.7, -0.5), ma = c(-1, -0.8)), n = n, sd = 1)
X[6,] <- x

## Generate y_t
A <- matrix(runif(p*p, -3, 3), ncol = p)
Y <- A*%X
Y <- t(Y)

## permutation = "max" or permutation = "fdr"
res <- PCA_TS(Y, lag.k = 5,permutation = "max")
res1 <- PCA_TS(Y, lag.k = 5,permutation = "fdr", beta = 10^(-10))
Z <- res$X

# Example 2 (Example 2 in the supplementary material of Chang, Guo and Yao (2018)).
# p=20, x_t consists of 5 independent subseries with 6, 5, 4, 3 and 2 components.

## Generate x_t
p <- 20;n <- 3000
X <- mat.or.vec(p,n)
x <- arima.sim(model = list(ar = c(0.5, 0.3), ma = c(-0.9, 0.3, 1.2, 1.3)),
n.start = 500, n = n+5, sd = 1)
for(i in 1:6) X[i,] <- x[i:(n+i-1)]
```

```

x <- arima.sim(model = list(ar = c(-0.4, 0.5), ma = c(1, 0.8, 1.5, 1.8)),
n.start = 500, n = n+4, sd = 1)
for(i in 7:11) X[i,] <- x[(i-6):(n+i-7)]
x <- arima.sim(model = list(ar = c(0.85,-0.3), ma=c(1, 0.5, 1.2)),
n.start = 500, n = n+3,sd = 1)
for(i in 12:15) X[i,] <- x[(i-11):(n+i-12)]
x <- arima.sim(model = list(ar = c(0.8, -0.5),ma = c(1, 0.8, 1.8)),
n.start = 500, n = n+2,sd = 1)
for(i in 16:18) X[i,] <- x[(i-15):(n+i-16)]
x <- arima.sim(model = list(ar = c(-0.7, -0.5), ma = c(-1, -0.8)),
n.start = 500,n = n+1,sd = 1)
for(i in 19:20) X[i,] <- x[(i-18):(n+i-19)]

## Generate y_t
A <- matrix(runif(p*p, -3, 3), ncol =p)
Y <- A%%X
Y <- t(Y)

## permutation = "max" or permutation = "fdr"
res <- PCA_TS(Y, lag.k = 5,permutation = "max")
res1 <- PCA_TS(Y, lag.k = 5,permutation = "fdr",beta = 10^(-200))
Z <- res$X

```

predict.factors

Make predictions from a "factors" object

Description

This function makes predictions from a "factors" object.

Usage

```

## S3 method for class 'factors'
predict(
  object,
  newdata = NULL,
  n.ahead = 10,
  control_ARIMA = list(),
  control_VAR = list(),
  ...
)

```

Arguments

object	An object of class "factors" constructed by Factors .
newdata	Optional. A new data matrix to predict from.
n.ahead	An integer specifying the number of steps ahead for prediction.

control_ARIMA	A list of arguments passed to the function <code>auto.arima()</code> of forecast . See 'Details' and the manual of <code>auto.arima()</code> . The default is <code>list(ic = "aic")</code> .
control_VAR	A list of arguments passed to the function <code>VAR()</code> of vars . See 'Details' and the manual of <code>VAR()</code> . The default is <code>list(type = "const", lag.max = 6, ic = "AIC")</code> .
...	Currently not used.

Details

Forecasting for \mathbf{y}_t can be implemented in two steps:

Step 1. Get the h -step ahead forecast of the $\hat{r} \times 1$ time series $\hat{\mathbf{x}}_t$ [See [Factors](#)], denoted by $\hat{\mathbf{x}}_{n+h}$, using a VAR model (if $\hat{r} > 1$) or an ARIMA model (if $\hat{r} = 1$). The orders of VAR and ARIMA models are determined by AIC by default. Otherwise, they can also be specified by users through the arguments `control_VAR` and `control_ARIMA`, respectively.

Step 2. The forecasted value for \mathbf{y}_t is obtained by $\hat{\mathbf{y}}_{n+h} = \hat{\mathbf{A}}\hat{\mathbf{x}}_{n+h}$.

Value

`ts_pred` A matrix of predicted values.

See Also

[Factors](#)

Examples

```
library(HDTSA)
data(FamaFrench, package = "HDTSA")

## Remove the market effects
reg <- lm(as.matrix(FamaFrench[, -c(1:2)]) ~ as.matrix(FamaFrench$MKT.RF))
Y_2d = reg$residuals

res_factors <- Factors(Y_2d, lag.k = 5)
pred_fac_Y <- predict(res_factors, n.ahead = 1)
```

predict.mtscp

Make predictions from a "mtscp" object

Description

This function makes predictions from a "mtscp" object.

Usage

```
## S3 method for class 'mtscp'
predict(
  object,
  newdata = NULL,
  n.ahead = 10,
  control_ARIMA = list(),
  control_VAR = list(),
  ...
)
```

Arguments

object	An object of class "mtscp" constructed by CP_MTS .
newdata	Optional. A new data matrix to predict from.
n.ahead	An integer specifying the number of steps ahead for prediction.
control_ARIMA	A list of arguments passed to the function <code>auto.arima()</code> of forecast . See 'Details' and the manual of <code>auto.arima()</code> . The default is <code>list(ic = "aic")</code> .
control_VAR	A list of arguments passed to the function <code>VAR()</code> of vars . See 'Details' and the manual of <code>VAR()</code> . The default is <code>list(type = "const", lag.max = 6, ic = "AIC")</code> .
...	Currently not used.

Details

Forecasting for \mathbf{y}_t can be implemented in two steps:

Step 1. Get the h -step ahead forecast of the $\hat{d} \times 1$ time series $\hat{\mathbf{x}}_t = (\hat{x}_{t,1}, \dots, \hat{x}_{t,\hat{d}})'$ [See [CP_MTS](#)], denoted by $\hat{\mathbf{x}}_{n+h}$, using a VAR model (if $\hat{d} > 1$) or an ARIMA model (if $\hat{d} = 1$). The orders of VAR and ARIMA models are determined by AIC by default. Otherwise, they can also be specified by users through the arguments `control_VAR` and `control_ARIMA`, respectively.

Step 2. The forecasted value for \mathbf{Y}_t is obtained by $\hat{\mathbf{Y}}_{n+h} = \hat{\mathbf{A}}\hat{\mathbf{X}}_{n+h}\hat{\mathbf{B}}'$ with $\hat{\mathbf{X}}_{n+h} = \text{diag}(\hat{\mathbf{x}}_{n+h})$.

Value

Y_pred	A list of length <code>n.ahead</code> , where each element is a $p \times q$ matrix representing the predicted values at each time step.
--------	--

See Also

[CP_MTS](#)

Examples

```
library(HDTSA)
data(FamaFrench, package = "HDTSA")

## Remove the market effects
```

```

reg <- lm(as.matrix(FamaFrench[, -c(1:2)]) ~ as.matrix(FamaFrench$MKT.RF))
Y_2d = reg$residuals

## Rearrange Y_2d into a 3-dimensional array Y
Y = array(NA, dim = c(NROW(Y_2d), 10, 10))
for (tt in 1:NROW(Y_2d)) {
  for (ii in 1:10) {
    Y[tt, ii, ] <- Y_2d[tt, (1 + 10*(ii - 1)):(10 * ii)]
  }
}

res_cp <- CP_MTS(Y, lag.k = 20, method = "CP.Refined")
pred_cp_Y <- predict(res_cp, n.ahead = 1)[[1]]

```

predict.tspca

Make predictions from a "tspca" object

Description

This function makes predictions from a "tspca" object.

Usage

```

## S3 method for class 'tspca'
predict(
  object,
  newdata = NULL,
  n.ahead = 10,
  control_ARIMA = list(),
  control_VAR = list(),
  ...
)

```

Arguments

object	An object of class "tspca" constructed by PCA_TS .
newdata	Optional. A new data matrix to predict from.
n.ahead	An integer specifying the number of steps ahead for prediction.
control_ARIMA	A list of arguments passed to the function <code>auto.arima()</code> of forecast . See 'Details' and the manual of <code>auto.arima()</code> . The default is <code>list(max.d = 0, max.q = 0, ic = "aic")</code> .
control_VAR	A list of arguments passed to the function <code>VAR()</code> of vars . See 'Details' and the manual of <code>VAR()</code> . The default is <code>list(type = "const", lag.max = 6, ic = "AIC")</code> .
...	Currently not used.

Details

Having obtained $\hat{\mathbf{x}}_t$ using the `PCA_TS` function, which is segmented into q uncorrelated subseries $\hat{\mathbf{x}}_t^{(1)}, \dots, \hat{\mathbf{x}}_t^{(q)}$, the forecasting for \mathbf{y}_t can be performed in two steps:

Step 1. Get the h -step ahead forecast $\hat{\mathbf{x}}_{n+h}^{(j)}$ ($j = 1, \dots, q$) by using a VAR model (if the dimension of $\hat{\mathbf{x}}_t^{(j)}$ is larger than 1) or an ARIMA model (if the dimension of $\hat{\mathbf{x}}_t^{(j)}$ is 1). The orders of VAR and ARIMA models are determined by AIC by default. Otherwise, they can also be specified by users through the arguments `control_VAR` and `control_ARIMA`, respectively.

Step 2. Let $\hat{\mathbf{x}}_{n+h} = (\{\hat{\mathbf{x}}_{n+h}^{(1)}\}', \dots, \{\hat{\mathbf{x}}_{n+h}^{(q)}\}')'$. The forecasted value for \mathbf{y}_t is obtained by $\hat{\mathbf{y}}_{n+h} = \hat{\mathbf{B}}^{-1}\hat{\mathbf{x}}_{n+h}$.

Value

`Y_pred` A matrix of predicted values.

See Also

[PCA_TS](#)

Examples

```
library(HDTSA)
data(FamaFrench, package = "HDTSA")

## Remove the market effects
reg <- lm(as.matrix(FamaFrench[, -c(1:2)]) ~ as.matrix(FamaFrench$MKT.RF))
Y_2d = reg$residuals

res_pca <- PCA_TS(Y_2d, lag.k = 5, thresh = TRUE)
pred_pca_Y <- predict(res_pca, n.ahead = 1)
```

QWIdata

The national QWI hires data

Description

The data on new hires at a national level are obtained from the Quarterly Workforce Indicators (QWI) of the Longitudinal Employer-Household Dynamics program at the U.S. Census Bureau (Abowd et al., 2009). The national QWI hires data covers a variable number of years, with some states providing time series going back to 1990 (e.g., Washington), and others (e.g., Massachusetts) only commencing at 2010. For each of 51 states (excluding D.C. but including Puerto Rico) there is a new hires time series for each county. Additional description of the data, along with its relevancy to labor economics, can be found in Hyatt and McElroy (2019).

Usage

```
data(QWIData)
```

Format

A list with 51 elements. Every element contains a multivariate time series.

Source

<https://qwiexplorer.ces.census.gov/>

<https://ledextract.ces.census.gov/qwi/all>

References

Abowd, J. M., Stephens, B. E., Vilhuber, L., Andersson, F., McKinney, K. L., Roemer, M., and Woodcock, S. (2009). The LEHD infrastructure files and the creation of the quarterly workforce indicators. In *Producer dynamics: New evidence from micro data*, pages 149–230. University of Chicago Press. doi:10.7208/chicago/9780226172576.003.0006.

Hyatt, H. R. and McElroy, T. S. (2019). Labor reallocation, employment, and earnings: Vector autoregression evidence. *Labour*, **33**, 463–487. doi:10.1111/labr.12153

SpecMulTest

Multiple testing with FDR control for spectral density matrix

Description

SpecMulTest() implements a new multiple testing procedure proposed in Chang et al. (2022) for the following Q hypothesis testing problems:

$$H_{0,q} : f_{i,j}(\omega) = 0 \text{ for any } (i, j) \in \mathcal{I}^{(q)} \text{ and } \omega \in \mathcal{J}^{(q)} \text{ versus } H_{1,q} : H_{0,q} \text{ is not true}$$

for $q = 1, \dots, Q$. Here, $f_{i,j}(\omega)$ represents the cross-spectral density between $x_{t,i}$ and $x_{t,j}$ at frequency ω with $x_{t,i}$ being the i -th component of the $p \times 1$ times series \mathbf{x}_t , and $\mathcal{I}^{(q)}$ and $\mathcal{J}^{(q)}$ refer to the set of index pairs and the set of frequencies associated with the q -th test, respectively.

Usage

```
SpecMulTest(Q, PVal, alpha = 0.05, seq_len = 0.01)
```

Arguments

Q	The number of the hypothesis tests.
PVal	A vector of length Q representing p-values of the Q hypothesis tests.
alpha	The prescribed level for the FDR control. The default is 0.05.
seq_len	The step size for generating a sequence from 0 to $\sqrt{2 \times \log Q - 2 \times \log(\log Q)}$. The default is 0.01.

Value

An object of class "hdtstest", which contains the following component:

MultiTest A logical vector of length Q . If its q -th element is TRUE, it indicates that $H_{0,q}$ should be rejected. Otherwise, $H_{0,q}$ should not be rejected.

References

Chang, J., Jiang, Q., McElroy, T. S., & Shao, X. (2022). Statistical inference for high-dimensional spectral density matrix. *arXiv preprint*. doi:10.48550/arXiv.2212.13686.

See Also

[SpecTest](#)

Examples

```
# Example 1
## Generate xt
n <- 200
p <- 10
flag_c <- 0.8
B <- 1000
burn <- 1000
z.sim <- matrix(rnorm((n+burn)*p),p,n+burn)
phi.mat <- 0.4*diag(p)
x.sim <- phi.mat %*% z.sim[, (burn+1):(burn+n)]
x <- x.sim - rowMeans(x.sim)
Q <- 4

## Generate the sets Iq and Jq
ISET <- list()
ISET[[1]] <- matrix(c(1,2),ncol=2)
ISET[[2]] <- matrix(c(1,3),ncol=2)
ISET[[3]] <- matrix(c(1,4),ncol=2)
ISET[[4]] <- matrix(c(1,5),ncol=2)
JSET <- as.list(2*pi*seq(0,3)/4 - pi)

## Calculate Q p-values
PVal <- rep(NA,Q)
for (q in 1:Q) {
  cross.indices <- ISET[[q]]
  J.set <- JSET[[q]]
  temp.q <- SpecTest(t(x), J.set, cross.indices, B, flag_c)
  PVal[q] <- temp.q$p.value
}
res <- SpecMulTest(Q, PVal)
res
```

Description

SpecTest() implements a new global test proposed in Chang et al. (2022) for the following hypothesis testing problem:

$$H_0 : f_{i,j}(\omega) = 0 \text{ for any } (i, j) \in \mathcal{I} \text{ and } \omega \in \mathcal{J} \text{ versus } H_1 : H_0 \text{ is not true,}$$

where $f_{i,j}(\omega)$ represents the cross-spectral density between $x_{t,i}$ and $x_{t,j}$ at frequency ω with $x_{t,i}$ being the i -th component of the $p \times 1$ times series \mathbf{x}_t . Here, \mathcal{I} is the set of index pairs of interest, and \mathcal{J} is the set of frequencies.

Usage

```
SpecTest(X, J.set, cross.indices, B = 1000, flag_c = 0.8)
```

Arguments

X	An $n \times p$ data matrix $\mathbf{X} = (\mathbf{x}_1, \dots, \mathbf{x}_n)'$, where n is the number of observations of the $p \times 1$ time series $\{\mathbf{x}_t\}_{t=1}^n$.
J.set	A vector representing the set \mathcal{J} of frequencies.
cross.indices	An $r \times 2$ matrix representing the set \mathcal{I} of r index pairs, where each row represents an index pair.
B	The number of bootstrap replications for generating multivariate normally distributed random vectors when calculating the critical value. The default is 2000.
flag_c	The bandwidth $c \in (0, 1]$ of the flat-top kernel for estimating $f_{i,j}(\omega)$ [See (2) in Chang et al. (2022)]. The default is 0.8.

Value

An object of class "hdtstest", which contains the following components:

Stat	The test statistic of the test.
pval	The p-value of the test.
cri95	The critical value of the test at the significance level 0.05.

References

Chang, J., Jiang, Q., McElroy, T. S., & Shao, X. (2022). Statistical inference for high-dimensional spectral density matrix. *arXiv preprint*. doi:10.48550/arXiv.2212.13686.

See Also

[SpecMulTest](#)

Examples

```

# Example 1
## Generate xt
n <- 200
p <- 10
flag_c <- 0.8
B <- 1000
burn <- 1000
z.sim <- matrix(rnorm((n+burn)*p),p,n+burn)
phi.mat <- 0.4*diag(p)
x.sim <- phi.mat %*% z.sim[, (burn+1):(burn+n)]
x <- x.sim - rowMeans(x.sim)

## Generate the sets I and J
cross.indices <- matrix(c(1,2), ncol=2)
J.set <- 2*pi*seq(0,3)/4 - pi

res <- SpecTest(t(x), J.set, cross.indices, B, flag_c)
Stat <- res$statistic
Pvalue <- res$p.value
CriVal <- res$cri95

```

UR_test

*Testing for unit roots based on sample autocovariances***Description**

This function implements the test proposed in Chang, Cheng and Yao (2022) for the following hypothesis testing problem:

$$H_0 : Y_t \sim I(0) \text{ versus } H_1 : Y_t \sim I(d) \text{ for some integer } d \geq 1,$$

where Y_t is a univariate time series.

Usage

```
UR_test(Y, lagk.vec = NULL, con_vec = NULL, alpha = 0.05)
```

Arguments

Y	A vector $\mathbf{Y} = (Y_1, \dots, Y_n)'$, where n is the number of the observations.
lagk.vec	The time lag K_0 used to calculate the test statistic [See Section 2.1 of Chang, Cheng and Yao (2022)]. It can be a vector specifying multiple time lags. If provided as a $s \times 1$ vector, the function will output the test results corresponding to each of the s values in lagk.vec. The default is $c(0, 1, 2, 3, 4)$.
con_vec	The constant c_κ specified in (5) of Chang, Cheng and Yao (2022). The default is 0.55. Alternatively, it can be an $m \times 1$ vector specified by users, representing m candidate values of c_κ .
alpha	The significance level of the test. The default is 0.05.

Value

An object of class "urtest", which contains the following components:

statistic	A $s \times 1$ vector with each element representing the test statistic value associated with each of the s time lags specified in lagk.vec.
reject	An $m \times s$ data matrix $\mathbf{R} = (R_{i,j})$ where $R_{i,j}$ represents whether the null hypothesis H_0 should be rejected for c_{κ} specified by the i -th component of con_vec, and K_0 specified by the j -th component of lagk.vec. $R_{i,j} = 1$ indicates rejection of the null hypothesis, while $R_{i,j} = 0$ indicates non-rejection.
lagk.vec	The time lags used in function.

References

Chang, J., Cheng, G., & Yao, Q. (2022). Testing for unit roots based on sample autocovariances. *Biometrika*, **109**, 543–550. doi:10.1093/biomet/asab034.

Examples

```
# Example 1
## Generate yt
N <- 100
Y <- arima.sim(list(ar = c(0.9)), n = 2*N, sd = sqrt(1))
con_vec <- c(0.45, 0.55, 0.65)
lagk.vec <- c(0, 1, 2)

UR_test(Y, lagk.vec = lagk.vec, con_vec = con_vec, alpha = 0.05)
UR_test(Y, alpha = 0.05)
```

 WN_test

Testing for white noise hypothesis in high dimension

Description

WN_test() implements the test proposed in Chang, Yao and Zhou (2017) for the following hypothesis testing problem:

$$H_0 : \{\mathbf{y}_t\}_{t=1}^n \text{ is white noise versus } H_1 : \{\mathbf{y}_t\}_{t=1}^n \text{ is not white noise.}$$

Usage

```
WN_test(
  Y,
  lag.k = 2,
  B = 1000,
  kernel.type = c("QS", "Par", "Bart"),
  pre = FALSE,
  alpha = 0.05,
  control.PCA = list()
)
```

Arguments

<code>Y</code>	An $n \times p$ data matrix $\mathbf{Y} = (\mathbf{y}_1, \dots, \mathbf{y}_n)'$, where n is the number of the observations of the $p \times 1$ time series $\{\mathbf{y}_t\}_{t=1}^n$.
<code>lag.k</code>	The time lag K used to calculate the test statistic [See (4) of Chang, Yao and Zhou (2017)]. The default is 2.
<code>B</code>	The number of bootstrap replications for generating multivariate normally distributed random vectors when calculating the critical value. The default is 1000.
<code>kernel.type</code>	The option for choosing the symmetric kernel used in the estimation of long-run covariance matrix. Available options include: "QS" (the default) for the Quadratic spectral kernel, "Par" for the Parzen kernel, and "Bart" for the Bartlett kernel. See Chang, Yao and Zhou (2017) for more information.
<code>pre</code>	Logical. If TRUE (the default), the time series PCA proposed in Chang, Guo and Yao (2018) should be performed on $\{\mathbf{y}_t\}_{t=1}^n$ before implementing the white noise test [See Remark 1 of Chang, Yao and Zhou (2017)]. The time series PCA is implemented by using the function <code>PCA_TS</code> with the arguments passed by <code>control.PCA</code> .
<code>alpha</code>	The significance level of the test. The default is 0.05.
<code>control.PCA</code>	A list of control arguments passed to the function <code>PCA_TS()</code> , including <code>lag.k</code> , <code>opt</code> , <code>thresh</code> , <code>delta</code> , and the associated arguments passed to the <code>clime</code> function (when <code>opt = 2</code>). See 'Details' in <code>PCA_TS</code> .

Value

An object of class "hdtstest", which contains the following components:

<code>statistic</code>	The test statistic of the test.
<code>p.value</code>	The p-value of the test.
<code>lag.k</code>	The time lag used in function.
<code>kernel.type</code>	The kernel used in function.

References

Chang, J., Guo, B., & Yao, Q. (2018). Principal component analysis for second-order stationary vector time series. *The Annals of Statistics*, **46**, 2094–2124. doi:10.1214/17AOS1613.

Chang, J., Yao, Q., & Zhou, W. (2017). Testing for high-dimensional white noise using maximum cross-correlations. *Biometrika*, **104**, 111–127. doi:10.1093/biomet/asw066.

See Also

[PCA_TS](#)

Examples

```
#Example 1
## Generate xt
n <- 200
p <- 10
Y <- matrix(rnorm(n * p), n, p)

res <- WN_test(Y)
Pvalue <- res$p.value
rej <- res$reject
```

Index

* **data**

FamaFrench, 11
IPindices, 14
QWIdata, 24

Coint, 2
CP_MTS, 5, 9, 22

DGP.CP, 8

Factors, 9, 13, 20, 21
FamaFrench, 11

HDSReg, 12

IPindices, 14

MartG_test, 15

PCA_TS, 17, 23, 24, 30
predict.factors, 20
predict.mtscp, 21
predict.tspca, 23

QWIdata, 24

SpecMulTest, 25, 27
SpecTest, 26, 27

UR_test, 28

WN_test, 29