Package 'FastBandChol'

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FastBandChol-package Fast estimation of covariance matrix by banded Cholesky factor

Description

Fast and numerically stable estimation of covariance matrix by banding the Cholesky factor using a modified Gram-Schmidt algorithm implemented in RcppArmadilo. See https://stat.umn.edu/~molst029 for details on the algorithm.

Details

Package: FastBandChol
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Author(s)

Aaron Molstad

References

Rothman, A.J., Levina, E., and Zhu, J. (2010). A new approach to Cholesky-based covariance regularization in high dimensions. Biometrika, 97(3):539-550.

```
## set sample size and dimension
n = 20
p = 100

## create covariance with AR1 structure
Sigma = matrix(0, nrow=p, ncol=p)
for(l in 1:p){
    for(m in 1:p){
        Sigma[l,m] = .5^(abs(l-m))
    }
}

## simulation Normal data
eo1 = eigen(Sigma)
Sigma.sqrt = eo1$vec%*%diag(eo1$val^.5)%*%t(eo1$vec)
X = t(Sigma.sqrt%*%matrix(rnorm(n*p), nrow=p, ncol=n))
```

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```
## compute estimates
est.sample = banded.sample(X, bandwidth=4)$est
est.chol = banded.chol(X, bandwidth=4)$est
```

banded.chol Computes estimate of covariance matrix by banding the Cholesky factor

Description

Computes estimate of covariance matrix by banding the Cholesky factor using a modified Gram Schmidt algorithm implemented in RcppArmadillo.

Usage

```
banded.chol(X, bandwidth, centered = FALSE)
```

Arguments

X A data matrix with n rows and p columns. Rows are assumed to be independent

realizations from a p-variate distribution with covariance Σ .

bandwidth A positive integer. Must be less than n-1 and p-1.

centered Logical. Is data matrix centered? Default is centered = FALSE

Value

A list with

est The estimated covariance matrix.

```
## set sample size and dimension
n=20
p=100

## create covariance with AR1 structure
Sigma = matrix(0, nrow=p, ncol=p)
for(l in 1:p){
    for(m in 1:p){
        Sigma[l,m] = .5^(abs(l-m))
    }
}

## simulation Normal data
eo1 = eigen(Sigma)
Sigma.sqrt = eo1$vec%*%diag(eo1$val^.5)%*%t(eo1$vec)
```

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```
X = t(Sigma.sqrt%*%matrix(rnorm(n*p), nrow=p, ncol=n))
## compute estimate
out1 = banded.chol(X, bandwidth=4)
```

banded.chol.cv

Selects bandwidth for Cholesky factorization by cross validation

Description

Selects bandwidth for Cholesky factorization by k-fold cross validation

Usage

```
banded.chol.cv(X, bandwidth, folds = 3, est.eval = TRUE, Frob = TRUE)
```

Arguments

X	A data matrix with n rows and p columns. Rows are assumed to be independent realizations from a p -variate distribution with covariance Σ .
bandwidth	A vector of candidate bandwidths. Candidate bandwidths can only positive integers such that the maximum is less than the sample size outside of the k th fold.
folds	The number of folds used for cross validation. Default is folds =3.
est.eval	Logical: est.eval = TRUE returns a list with both the selected bandwidth and the estimated covariance matrix. est.eval=FALSE returns a list with only the

selected bandwidth. The default is est.eval = TRUE.

Logical: Frob = TRUE uses squared Frobenius norm loss for cross-validation.

Frob = FALSE uses operator norm loss. Default is Frob = TRUE.

Value

a list with

Frob

bandwidth.min The bandwidth minimizing cross-validation error.

est The estimated covariance matrix computed with bandwidth=bandwidth.min.

```
## set sample size and dimension
n=20
p=100

## create covariance with AR1 structure
Sigma = matrix(0, nrow=p, ncol=p)
for(l in 1:p){
  for(m in 1:p){
```

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```
Sigma[1,m] = .5^(abs(1-m))
}

## simulation Normal data
eo1 = eigen(Sigma)
Sigma.sqrt = eo1$vec%*%diag(eo1$val^.5)%*%t(eo1$vec)
X = t(Sigma.sqrt%*%matrix(rnorm(n*p), nrow=p, ncol=n))

## perform cross validation
k = 4:7
out1.cv = banded.chol.cv(X, bandwidth=k, folds = 5)
```

banded.sample

Computes banded sample covariance matrix

Description

Estimates a covariance matrix by banding the sample covariance matrix

Usage

```
banded.sample(X, bandwidth, centered = FALSE)
```

Arguments

X A data matrix with n rows and p columns. Rows are assumed to be independent

realizations from a p-variate distribution with covariance Σ .

bandwidth A positive integer. Must be less than p-1..

centered Logical. Is data matrix centered? Default is centered = FALSE

Value

A list with

est The estimated covariance matrix.

```
## set sample size and dimension
n=20
p=100

## create covariance with AR1 structure
Sigma = matrix(0, nrow=p, ncol=p)
for(l in 1:p){
   for(m in 1:p){
      Sigma[l,m] = .5^(abs(l-m))
```

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```
}
}

## simulation Normal data
eo1 = eigen(Sigma)
Sigma.sqrt = eo1$vec%*%diag(eo1$val^.5)%*%t(eo1$vec)
X = t(Sigma.sqrt%*matrix(rnorm(n*p), nrow=p, ncol=n))
## compute estimate
out2 = banded.sample(X, bandwidth=4)
```

banded.sample.cv

Selects bandwidth for sample covariance matrix by cross validation

Description

Selects bandwidth for sample covariance matrix by k-fold cross validation

Usage

```
banded.sample.cv(X, bandwidth, folds = 3, est.eval = TRUE, Frob = TRUE)
```

Arguments

Χ	A data matrix with n rows and p columns. Rows are assumed to be independent realizations from a p -variate distribution with covariance Σ .
bandwidth	A vector of candidate bandwidths. Candidate bandwidths can only positive integers such that the maximum is less than $p-1$.
folds	The number of folds used for cross validation. Default is folds =3.
est.eval	Logical: est.eval = TRUE returns a list with both the selected bandwidth and the estimated covariance matrix. est.eval=FALSE returns a list with only the selected bandwidth. The default is est.eval = TRUE.
Frob	Logical: Frob = TRUE uses squared Frobenius norm loss for cross-validation. Frob = FALSE uses operator norm loss. Default is Frob = TRUE.

Value

A list with

bandwidth.min the bandwidth minimizing cv error
est the sample covariance matrix at bandwidth.min

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```
## set sample size and dimension
n=20
p=100
## create covariance with AR1 structure
Sigma = matrix(0, nrow=p, ncol=p)
for(l in 1:p){
  for(m in 1:p){
    Sigma[1,m] = .5^(abs(1-m))
  }
}
## simulation Normal data
eo1 = eigen(Sigma)
Sigma.sqrt = eo1$vec%*%diag(eo1$val^.5)%*%t(eo1$vec)
X = t(Sigma.sqrt%*%matrix(rnorm(n*p), nrow=p, ncol=n))
## perform cross validation
k = 4:7
out2.cv = banded.sample.cv(X, bandwidth=k, folds=5)
```

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