# Package 'HDTD'

October 17, 2020

Type Package

**Title** Statistical Inference about the Mean Matrix and the Covariance Matrices in High-Dimensional Transposable Data (HDTD)

Version 1.22.0

**Depends** R (>= 3.6)

Suggests knitr, markdown

**Imports** stats, Rcpp (>= 1.0.1)

LinkingTo Rcpp, RcppArmadillo

### **Description**

Characterization of intra-individual variability using physiologically relevant measurements provides important insights into fundamental biological questions ranging from cell type identity to tumor development. For each individual, the data measurements can be written as a matrix with the different subsamples of the individual recorded in the columns and the different phenotypic units recorded in the rows. Datasets of this type are called high-dimensional transposable data. The HDTD package provides functions for conducting statistical inference for the mean relationship between the row and column variables and for the covariance structure within and between the row and column variables.

License GPL-3

**biocViews** DifferentialExpression, Genetics, GeneExpression, Microarray, Sequencing, StatisticalMethod, Software

VignetteBuilder knitr

URL http://github.com/AnestisTouloumis/HDTD

BugReports http://github.com/AnestisTouloumis/HDTD/issues

LazyData true

NeedsCompilation no

RoxygenNote 7.0.2

**Encoding** UTF-8

git\_url https://git.bioconductor.org/packages/HDTD

git\_branch RELEASE\_3\_11

git\_last\_commit 984ac7b

git\_last\_commit\_date 2020-04-27

**Date/Publication** 2020-10-16

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## R topics documented:

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## **Description**

The package HDTD offers functions to estimate and test the matrix parameters of transposable data in high-dimensional settings.

## **Details**

The term transposable data refers to datasets that are structured in a matrix form such that both the rows and columns correspond to variables of interest. For example, consider microarray studies in genetics where multiple RNA samples across different tissues are available per subject. In this case, a data matrix can be created with row variables the genes, column variables the tissues and measurements the corresponding expression levels.

The function meanmat.hat estimates the mean matrix of the transposable data.

The mean relationship of the row and column variables can be tested using the function meanmat.ts. The implemented test is nonparametric and not seriously restricted by the dependence structure among and/or between the row and column variables. See *Touloumis et al.* (2015) for more details.

The function covmat.hat provides Stein-type shrinkage estimators for the row covariance matrix and/or for the column covariance matrix under a matrix-variate normal model. See *Touloumis et al.* (2016) for more details.

The sphericity and identity hypothesis for the row or column covariance matrix can be tested using the function covmat.ts. Both tests are nonparametric, i.e., they do not rely on a normality assumption. See *Touloumis et al.* (2017) for more details.

There are three utility functions that allow the user to change to interchange the role of row and column variables (transposedata), to center the transposable data (centerdata) or to rearrange the order of the row and/or column variables (orderdata).

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#### Author(s)

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#### References

Touloumis, A., Tavare, S. and Marioni, J. C. (2015) Testing the Mean Matrix in High-Dimensional Transposable Data. *Biometrics* **71**, 157–166

Touloumis, A., Marioni, J. C. and Tavare, S. (2016) HDTD: Analyzing multi-tissue gene expression data. *Bioinformatics* **32**, 2193–2195.

Touloumis, A., Marioni, J. C. and Tavare, S. (2019+) Hypothesis Testing for the Covariance Matrix in High-Dimensional Transposable Data with Kronecker Product Dependence Structure. *Statistica Sinica*.

## **Examples**

```
data(VEGFmouse)
## The sample mean matrix.
sample_mean <- meanmat.hat(datamat = VEGFmouse, N = 40)
sample_mean
## Testing conservation of the overall gene expression across tissues.
tissues_mean_test <- meanmat.ts(datamat = VEGFmouse, N = 40, group.sizes = 9)
tissues_mean_test
# Estimating the gene and column covariance matrices.
est_cov_mat <- covmat.hat(datamat = VEGFmouse, N = 40)
est_cov_mat
## Hypothesis tests for the covariance matrix of the genes (rows).
genes_cov_test <- covmat.ts(datamat = VEGFmouse, N = 40)
genes_cov_test
## Hypothesis tests for the covariance matrix of the tissues (columns).
tissues_cov_test <- covmat.ts(datamat = VEGFmouse, N = 40, voi = 'columns')
tissues_cov_test</pre>
```

centerdata

Centering Transposable Data

## **Description**

This function centers the transposable data around their sample mean matrix.

## Usage

```
centerdata(datamat, N)
```

## **Arguments**

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, e.g., the number of subjects.

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### **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

### Value

Returns a matrix of the same size as datamat.

#### Author(s)

Anestis Touloumis

### See Also

```
covmat.hat and covmat.ts.
```

## **Examples**

```
data(VEGFmouse)
## Centering the VEGF dataset around the sample mean matrix.
VEGFcen <- centerdata(datamat = VEGFmouse, N = 40)</pre>
```

covmat.hat

Estimation of the Row and of the Column Covariance Matrices.

## Description

This function provides the row and/or column covariance matrix estimators.

## Usage

```
covmat.hat(datamat, N, shrink = "both", centered = FALSE, voi = "both")
```

## Arguments

datamat	numeric matrix containing the transposable data.
N	positive integer number indicating the sample size, i.e., the number of subjects.
shrink	character indicating if shrinkage estimation should be performed. Options include 'rows', 'columns', 'both' and 'none'.
centered	logical indicating if the transposable data are centered. Options include TRUE or FALSE.
voi	character indicating if the row, column or both covariance matrices should be printed. Options include 'rows', 'columns' and 'both'.

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#### **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

For identifiability reasons, the trace of the row covariance matrix is set equal to its dimension. If you want to place the equivalent restriction on the column covariance matrix, interchange the role of row and column variables by utilizing the function transposedata.

#### Value

Returns a list with components:

rows.covmat the estimated row covariance matrix.

rows.intensity the estimated row intensity.

cols.covmat the estimated column covariance matrix.

cols.intensity the estimated column intensity.

N the sample size.

n.rows the number of row variables.n.cols the number of column variables.

shrink character indicating if shrinkage estimation was performed. centered logical indicating if the transposable data were centered.

### Author(s)

**Anestis Touloumis** 

## References

Touloumis, A., Marioni, J. C. and Tavare, S. (2016) HDTD: Analyzing multi-tissue gene expression data, *Bioinformatics* **32**, 2193–2195.

## **Examples**

```
data(VEGFmouse)
# Estimating the gene and tissue covariance matrices.
est_cov_mat <- covmat.hat(datamat = VEGFmouse, N = 40)
est_cov_mat</pre>
```

covmat.ts

Nonparametric Tests for the Row or Column Covariance Matrix

## **Description**

Testing the sphericity, identity and diagonality hypotheses for the row or column covariance matrix.

## Usage

```
covmat.ts(datamat = datamat, N = N, voi = "rows", centered = FALSE)
```

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## Arguments

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, i.e., the number of subjects. voi character indicating if the test should be applied on the row or column covariance

matrix. Options include 'rows' or 'columns'.

centered logical indicating if the transposable data are centered. Options include TRUE or

FALSE.

#### **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

The tests are nonparametric and thus robust to some departures from the matrix-variate normal model.

#### Value

It returns a list with components:

diagonality.ts a list containing the test statistic and p-value of the diagonality hypothesis test. sphericity.ts a list containing the test statistic and p-value of the sphericity hypothesis test.

identity.ts a list containing the test statistic and p-value of the identity hypothesis test.

N the sample size.

n.rows the number of row variables.n.cols the number of column variables.

variables character indicating if the tests were applied to the row or column covariance

matrix.

centered logical indicating if the transposable data were centered.

## Author(s)

Anestis Touloumis

### References

Touloumis, A., Marioni, J.C. and Tavare, S. (2019+). Hypothesis Testing for the Covariance Matrix in High-Dimensional Transposable Data with Kronecker Product Dependence Structure. *Statistica Sinica*.

```
data(VEGFmouse)
## Hypothesis tests for the covariance matrix of the genes (rows).
genes_cov_test <- covmat.ts(datamat = VEGFmouse, N = 40)
genes_cov_test
## Hypothesis tests for the covariance matrix of the tissues (columns).
tissues_cov_test <- covmat.ts(datamat = VEGFmouse, N = 40, voi = 'columns')
tissues_cov_test</pre>
```

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meanmat.hat Estimation the Mean Matrix
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## **Description**

This function estimates the mean matrix.

### Usage

```
meanmat.hat(datamat, N, group.sizes = NULL, group.vars = NULL)
```

### **Arguments**

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, i.e., the number of subjects. group.sizes numeric vector indicating the size of the row or column groups that share the

same mean vector. It should be used only when group  $.\,\mbox{vars='rows'}$  or 'columns'.

group.vars character indicating that the mean matrix can be simplified over the row or col-

umn variables. Options include 'rows' or 'columns'.

### **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

## Value

Returns a list with components:

estmeanmat the estimated mean matrix.

N the sample size.

n.rows the number of row variables.n.cols the number of column variables.

## Author(s)

**Anestis Touloumis** 

#### References

Touloumis, A., Marioni, J. C. and Tavare, S. (2016) HDTD: Analyzing multi-tissue gene expression data. *Bioinformatics* **32**, 2193–2195.

```
data(VEGFmouse)
## The sample mean matrix of the VEGF mouse data.
sample_mean <- meanmat.hat(datamat = VEGFmouse, N = 40)
sample_mean
sample_mean$estmeanmat</pre>
```

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meanmat.ts
------------

Nonparametric Tests for the Mean Matrix

### **Description**

This function performs hypothesis testing for the mean matrix.

## Usage

```
meanmat.ts(datamat, N, group.sizes, voi = "columns")
```

### **Arguments**

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, i.e., the number of subjects.

group.sizes numeric vector indicating the group sizes under the null hypothesis.

voi character indicating if the test will be applied to the row or column variables.

Options include 'rows' or 'columns'.

#### Details

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

## Value

Returns a list with components:

statistic the value of the test statistic.
p.value the corresponding p-value.

voi the set of variables that the test was applied to.

n.groups the number of groups under the null hypothesis.
group.sizes the size of each group under the null hypothesis.

N the sample size.

n.rows the number of row variables.n.cols the number of column variables.

## Author(s)

Anestis Touloumis

### References

Touloumis, A., Tavare, S. and Marioni, J. C. (2015) Testing the Mean Matrix in High-Dimensional Transposable Data. *Biometrics* **71**, 157–166.

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## **Examples**

```
data(VEGFmouse)
## Testing conservation of the overall gene expression across tissues.
tissues_mean_test <- meanmat.ts(datamat = VEGFmouse, N = 40, group.sizes = 9)
tissues_mean_test
## Testing if the adrenal and the cerebrum tissues have the same mean vector.
test2 <- meanmat.ts(VEGFmouse, N = 40, group.sizes = c(2, rep(1,7)))
test2</pre>
```

orderdata

Reordering Row and Column Variables

## **Description**

This utility function rearranges the row and/or the column variables in a desired order.

## Usage

```
orderdata(datamat, N, order.rows = NULL, order.cols = NULL)
```

## **Arguments**

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, i.e., the number of subjects.

order.rows numeric vector displaying the desired order of the row variables.

order.cols numeric vector displaying the desired order of the column variables.

## **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

## Value

Returns a matrix of the same size as datamat.

## Author(s)

Anestis Touloumis

## See Also

```
meanmat.ts and meanmat.hat.
```

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#### **Examples**

```
data(VEGFmouse)
set.seed(1)
tissuesold <- colnames(VEGFmouse[ ,1:9])
## Suppose that you want to order the tissues in the folowing order.
tissuesnew <- colnames(VEGFmouse[ ,1:9])[sample(9)]
tissuesnew
## To do this, create a numeric vector with the desired order.
ordtis <- pmatch(tissuesnew, tissuesold)
VEGFmousenew <- orderdata(datamat = VEGFmouse, N = 40, order.cols = ordtis)
colnames(VEGFmousenew)[1:9]</pre>
```

transposedata

Interchanging the Row and Column Variables in Transposable Data

## Description

This function interchanges the row and column variables in transposable data so that the original row variables will be treated as column variables and the original column variables as row variables.

## Usage

```
transposedata(datamat, N)
```

### **Arguments**

datamat numeric matrix containing the transposable data.

N positive integer number indicating the sample size, i.e., the number of subjects.

## **Details**

It is assumed that there are nrow(datamat) row variables and ncol(datamat)/N column variables in datamat. Further, datamat should be written in such a way that every ncol(datamat)/N consecutive columns belong to the same subject and the order of the column variables in each block is preserved across subjects.

#### Value

Returns a matrix with ncol(datamat) rows and nrow(datamat)/N columns.

## Author(s)

**Anestis Touloumis** 

#### See Also

centerdata and orderdata.

```
data(VEGFmouse)
## Transposing the VEGF dataset.
VEGFtr <- transposedata(datamat = VEGFmouse, N = 40)</pre>
```

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**VEGFmouse** 

Vascular Endothelial Growth Factor Mouse Dataset

## Description

Log2 normalized mouse gene expression data in the vascular endothelial growth factor signalling pathway across multiple tissues.

### **Format**

A data frame with 46 rows and 360 columns. The rows corresponds to 46 genes in the VEGF signalling pathway. The column names indicate the mouse and the tissue on which gene expression levels were measured. Since there are 40 mice and 9 tissues, we have a total of 360 columns. Every 9 consecutive columns belong to the same mouse and the tissues are ordered in the same way in each mouse.

## **Source**

Zahn et al. (2007). AGEMAP: A gene expression database for aging in mice. *PLoS Genetics* 3, e201.

```
data(VEGFmouse)
## Check the order of the tissues from the first mouse.
colnames(VEGFmouse[,1:9])
```

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