# Package 'HDNRA'

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```
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Title High-Dimensional Location Testing with Normal-Reference
           Approaches
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Description We provide a collection of various classical tests and latest normal-
           reference tests for comparing high-dimensional mean vectors including two-sample and gen-
           eral linear hypothesis testing (GLHT) problem. Some existing tests for two-sample prob-
           lem [see Bai, Zhidong, and Hewa Saranadasa.(1996) <a href="https://example.com/https:/
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           lem [see Zhang, Jin-
```

2 Contents

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## **Contents**

BS1996.TS.NABT
corneal
COVID19
CQ2010.TSBF.NABT
FHW2004.GLHT.NABT
NRtest.object
print.NRtest
S2007.ks.NABT
SD2008.TS.NABT
SF2006.GLHT.NABT
SKK2013.TSBF.NABT
YS2012.GLHT.NABT
ZGZ2017.GLHT.2cNRT
ZGZ2017.GLHTBF.NABT
ZGZC2020.TS.2cNRT
ZWZ2023.TSBF.2cNRT
ZZ2022.GLHT.3cNRT

BS1996.TS.NABT 3

BS19	96.TS.NABT	Norm by Ba							est	fo	r	tw	0-	sai	mį	əle	p	ro	— ble	 ?m	p	ro	ро	se	d
Index																									38
	ZZZ2023.TSBF.20	cNRT .	•				•		•	•		•				•			•		•			•	36
	ZZZ2022.GLHT.2	2cNRT				 																			34
	ZZZ2020.TS.2cN	RT				 																			33
	ZZGZ2021.TSBF.	2cNRT				 																			32
	ZZG2022.GLHTE	3F.2cNR	Γ.			 																			30
	ZZ2022.TSBF.3cl	NRT				 																			29
	ZZ2022.TS.3cNR	Т				 																			28
	ZZ2022.GLHTBF	3cNRT				 																			26

## Description

Bai and Saranadasa (1996)'s test for testing equality of two-sample high-dimensional mean vectors with assuming that two covariance matrices are the same.

### Usage

BS1996.TS.NABT(y1, y2)

### Arguments

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a $p$ -dimensional observation.
y2	The data matrix $(n_2 \times p)$ from the second population. Each row represents a $p$ -dimensional observation.

## **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1},\ldots,\boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1})=\boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1})=\boldsymbol{\Sigma}, i=1,2$ .

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Bai and Saranadasa (1996) proposed the following centralised  $L^2$ -norm-based test statistic:

$$T_{BS} = \frac{n_1 n_2}{n} \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2 - \operatorname{tr}(\hat{\boldsymbol{\Sigma}}),$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors and  $\hat{\Sigma}$  is the pooled sample covariance matrix. They showed that under the null hypothesis,  $T_{BS}$  is asymptotically normally distributed.

4 corneal

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Bai Z, Saranadasa H (1996). "Effect of high dimension: by an example of a two sample problem." *Statistica Sinica*, 311–329. https://www.jstor.org/stable/24306018.

### **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
BS1996.TS.NABT(group1,group2)</pre>
```

corneal

HDNRA\_data corneal

#### **Description**

This dataset was acquired during a keratoconus study, a collaborative project involving Ms.Nancy Tripoli and Dr.Kenneth L.Cohen of Department of Ophthalmology at the University of North Carolina, Chapel Hill. The fitted feature vectors for the complete corneal surface dataset collectively into a feature matrix with dimensions of  $150 \times 2000$ .

### Usage

```
data(corneal)
```

#### **Format**

#### 'corneal':

A data frame with 150 observations on the following 4 groups.

**normal group1** row 1 to row 43 in total 43 rows of the feature matrix correspond to observations from the normal group

unilateral suspect group2 row 44 to row 57 in total 14 rows of the feature matrix correspond to observations from the unilateral suspect group

**suspect map group3** row 58 to row 78 in total 21 of the feature matrix correspond to observations from the suspect map group

**clinical keratoconus group4** row 79 to row 150 in total 72 of the feature matrix correspond to observations from the clinical keratoconus group

COVID19 5

#### References

Smaga Ł, Zhang J (2019). "Linear hypothesis testing with functional data." *Technometrics*, **61**(1), 99–110. doi:10.1080/00401706.2018.1456976.

### **Examples**

```
library(HDNRA)
data(corneal)
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group
dim(group1)
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group
dim(group2)
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group
dim(group3)
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group
dim(group4)</pre>
```

COVID19

HDNRA\_data COVID19

## Description

A COVID19 data set from NCBI with ID GSE152641. The data set profiled peripheral blood from 24 healthy controls and 62 prospectively enrolled patients with community-acquired lower respiratory tract infection by SARS-COV-2 within the first 24 hours of hospital admission using RNA sequencing.

#### Usage

```
data(COVID19)
```

#### **Format**

## 'COVID19':

A data frame with 86 observations on the following 2 groups.

**healthy group1** row 2 to row 19, and row 82 to 87, in total 24 healthy controls **patients group2** row 20 to 81, in total 62 prospectively enrolled patients

## References

Thair SA, He YD, Hasin-Brumshtein Y, Sakaram S, Pandya R, Toh J, Rawling D, Remmel M, Coyle S, Dalekos GN, others (2021). "Transcriptomic similarities and differences in host response between SARS-CoV-2 and other viral infections." *Iscience*, **24**(1). doi:10.1016/j.isci.2020.101947.

### **Examples**

```
library(HDNRA)
data(COVID19)
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
dim(group1)
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
dim(group2)</pre>
```

CQ2010.TSBF.NABT

Normal-approximation-based test for two-sample BF problem proposed by Chen and Qin (2010)

### **Description**

Chen and Qin (2010)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

#### Usage

```
CQ2010.TSBF.NABT(y1, y2)
```

#### **Arguments**

y1 The data matrix  $(n_1 \times p)$  from the first population. Each row represents a p-dimensional observation.

y2 The data matrix  $(n_2 \times p)$  from the second population. Each row represents a p-dimensional observation.

## **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Chen and Qin (2010) proposed the following test statistic:

$$T_{CQ} = rac{\sum_{i 
eq j}^{n_1} oldsymbol{y}_{1i}^ op oldsymbol{y}_{1j}}{n_1(n_1-1)} + rac{\sum_{i 
eq j}^{n_2} oldsymbol{y}_{2i}^ op oldsymbol{y}_{2j}}{n_2(n_2-1)} - 2rac{\sum_{i=1}^{n_1} \sum_{j=1}^{n_2} oldsymbol{y}_{1i}^ op oldsymbol{y}_{2j}}{n_1n_2}.$$

They showed that under the null hypothesis,  $T_{CQ}$  is asymptotically normally distributed.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

FHW2004.GLHT.NABT 7

#### References

Chen SX, Qin Y (2010). "A two-sample test for high-dimensional data with applications to gene-set testing." *The Annals of Statistics*, **38**(2). doi:10.1214/09aos716.

## **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
CQ2010.TSBF.NABT(group1,group2)</pre>
```

FHW2004.GLHT.NABT

Normal-approximation-based test for GLHT problem proposed by Fujikoshi et al. (2004)

## Description

Fujikoshi et al. (2004)'s test for general linear hypothesis testing (GLHT) problem for highdimensional data with assuming that underlying covariance matrices are the same.

#### Usage

```
FHW2004.GLHT.NABT(Y,X,C,n,p)
```

## Arguments

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
Χ	A known $n \times k$ full-rank design matrix with rank $(\mathbf{X}) = k < n$ .
С	A known matrix of size $q \times k$ with rank $(C) = q < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

#### **Details**

A high-dimensional linear regression model can be expressed as

$$Y = X\Theta + \epsilon$$
,

where  $\Theta$  is a  $k \times p$  unknown parameter matrix and  $\epsilon$  is an  $n \times p$  error matrix.

8 FHW2004.GLHT.NABT

It is of interest to test the following GLHT problem

$$H_0: C\Theta = \mathbf{0}, \quad \text{vs.} \quad H_1: C\Theta \neq \mathbf{0}.$$

Fujikoshi et al. (2004) proposed the following test statistic:

$$T_{FHW} = \sqrt{p} \left[ (n-k) \frac{\mathrm{tr}(\boldsymbol{S}_h)}{\mathrm{tr}(\boldsymbol{S}_e)} - q \right],$$

where  $S_h$  and  $S_e$  are the matrices of sums of squares and products due to the hypothesis and the error, respecitively.

They showed that under the null hypothesis,  $T_{FHW}$  is asymptotically normally distributed.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Fujikoshi Y, Himeno T, Wakaki H (2004). "Asymptotic results of a high dimensional MANOVA test and power comparison when the dimension is large compared to the sample size." *Journal of the Japan Statistical Society*, **34**(1), 19–26. doi:10.14490/jjss.34.19.

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
X \leftarrow matrix(c(rep(1,n[1]),rep(0,sum(n)),rep(1,n[2]), rep(0,sum(n)),
             rep(1,n[3]),rep(0,sum(n)),rep(1,n[4])),ncol=k,nrow=sum(n))
q < - k-1
C <- cbind(diag(q),-rep(1,q))</pre>
FHW2004.GLHT.NABT(Y,X,C,n,p)
```

NRtest.object 9

NRtest.object

S3 Class "NRtest"

## **Description**

The "NRtest" objects provide a comprehensive summary of hypothesis test outcomes, including test statistics, p-values, parameter estimates, and confidence intervals, if applicable.

## Usage

```
NRtest.object(
    statistic,
    p.value,
    method,
    null.value,
    alternative,
    parameter = NULL,
    sample.size = NULL,
    sample.dimension = NULL,
    estimation.method = NULL,
    data.name = NULL,
    ...
)
```

### **Arguments**

statistic	Numeric scalar containing the value of the test statistic, with a names attribute indicating the name of the test statistic.
p.value	Numeric scalar containing the p-value for the test.
method	Character string giving the name of the test.
null.value	Character string indicating the null hypothesis.
alternative	Character string indicating the alternative hypothesis.
parameter	Numeric vector containing the estimated approximation parameter(s) associated with the approximation method. This vector has a names attribute describing its element(s).
sample.size	Numeric vector containing the number of observations in each group used for the hypothesis test.
sample.dimension	on
	Numeric scalar containing the dimension of the dataset used for the hypothesis test.
estimation.meth	nod
	Character string giving the name of the approximation approach used to approximate the null distribution of the test statistic.
data.name	Character string describing the data set used in the hypothesis test.

Additional optional arguments.

NRtest.object

#### **Details**

A class of objects returned by high-dimensional hypothesis testing functions in the **HDNRA** package, designed to encapsulate detailed results from statistical hypothesis tests. These objects are structured similarly to **htest** objects in the package **EnvStats** but are tailored to the needs of the **HDNRA** package.

#### Value

An object of class "NRtest" containing both required and optional components depending on the specifics of the hypothesis test, shown as follows:

### **Required Components**

```
These components must be present in every "NRtest" object:
```

```
p.value Must e present.

null.value Must e present.

alternative Must e present.

method Must e present.
```

### **Optional Components**

These components are included depending on the specifics of the hypothesis test performed:

```
parameter May be present.
sample.size May be present.
sample.dimension May be present.
estimation.method May be present.
data.name May be present.
```

#### Methods

The class has the following methods:

```
print.NRtest Printing the contents of the NRtest object in a human-readable form.
```

```
# Example 1: Using Bai and Saranadasa (1996)'s test (two-sample problem)
NRtest.obj1 <- NRtest.object(
    statistic = c("T[BS]" = 2.208),
    p.value = 0.0136,
    method = "Bai and Saranadasa (1996)'s test",
    data.name = "group1 and group2",
    null.value = c("Difference between two mean vectors is o"),
    alternative = "Difference between two mean vectors is not 0",
    parameter = NULL,</pre>
```

print.NRtest 11

```
sample.size = c(n1 = 24, n2 = 26),
 sample.dimension = 20460,
 estimation.method = "Normal approximation"
)
print(NRtest.obj1)
# Example 2: Using Fujikoshi et al. (2004)'s test (GLHT problem)
NRtest.obj2 <- NRtest.object(
 statistic = c("T[FHW]" = 6.4015),
 p.value = 0,
 method = "Fujikoshi et al. (2004)'s test",
 data.name = "Y",
 null.value = "The general linear hypothesis is true",
 alternative = "The general linear hypothesis is not true",
 parameter = NULL,
 sample.size = c(n1 = 43, n2 = 14, n3 = 21, n4 = 72),
 sample.dimension = 2000,
 estimation.method = "Normal approximation"
)
print(NRtest.obj2)
```

print.NRtest

Print Method for S3 Class "NRtest"

### **Description**

Prints the details of the **NRtest** object in a user-friendly manner. This method provides a clear and concise presentation of the test results contained within the **NRtest** object, including all relevant statistical metrics and test details.

## Usage

```
## S3 method for class \pkg{NRtest}
## S3 method for class 'NRtest'
print(x, ...)
```

### **Arguments**

x an **NRtest** object.

... further arguments passed to or from other methods.

### **Details**

The print.NRtest function formats and presents the contents of the **NRtest** object, which includes statistical test results and related parameters. This function is designed to provide a user-friendly display of the object's contents, making it easier to understand the results of the analysis.

12 S2007.ks.NABT

#### Value

Invisibly returns the input x.

#### Author(s)

Pengfei Wang <nie23.wp8738@e.ntu.edu.sg>

#### See Also

NRtest.object

S2007.ks.NABT

Normal-approximation-based test for one-way MANOVA problem proposed by Schott (2007)

### **Description**

Schott, J. R. (2007)'s test for one-way MANOVA problem for high-dimensional data with assuming that underlying covariance matrices are the same.

### Usage

S2007.ks.NABT(Y, n, p)

### **Arguments**

Υ	A list of k data matrices. The ith element represents the data matrix $(n_i \times p)$
	from the $i$ th population with each row representing a $p$ -dimensional observation.
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group
	$i, n_i$ .
р	The dimension of data.

### **Details**

Suppose we have the following k independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}, i = 1, \dots, k$ .

It is of interest to test the following one-way MANOVA problem:

$$H_0: \boldsymbol{\mu}_1 = \dots = \boldsymbol{\mu}_k, \quad \text{ vs. } H_1: H_0 \text{ is not ture }.$$

Schott (2007) proposed the following test statistic:

$$T_S = [\operatorname{tr}(\boldsymbol{H})/h - \operatorname{tr}(\boldsymbol{E})/e]/\sqrt{N-1},$$

where  $\boldsymbol{H} = \sum_{i=1}^k n_i (\bar{\boldsymbol{y}}_i - \bar{\boldsymbol{y}}) (\bar{\boldsymbol{y}}_i - \bar{\boldsymbol{y}})^{\top}$ ,  $\boldsymbol{E} = \sum_{i=1}^k \sum_{j=1}^{n_i} (\boldsymbol{y}_{ij} - \bar{\boldsymbol{y}}_i) (\boldsymbol{y}_{ij} - \bar{\boldsymbol{y}}_i)^{\top}$ , h = k-1, and e = N-k, with  $N = n_1 + \dots + n_k$ . They showed that under the null hypothesis,  $T_S$  is asymptotically normally distributed.

SD2008.TS.NABT 13

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Schott JR (2007). "Some high-dimensional tests for a one-way MANOVA." *Journal of Multivariate Analysis*, **98**(9), 1825–1839. doi:10.1016/j.jmva.2006.11.007.

## **Examples**

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group
p <- dim(corneal)[2]
Y <- list()
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))
S2007.ks.NABT(Y, n, p)</pre>
```

SD2008.TS.NABT

Normal-approximation-based test for two-sample problem proposed by Srivastava and Du (2008)

## **Description**

Srivastava and Du (2008)'s test for testing equality of two-sample high-dimensional mean vectors with assuming that two covariance matrices are the same.

## Usage

```
SD2008.TS.NABT(y1, y2)
```

### **Arguments**

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a p-
	dimensional observation.
y2	The data matrix $(n_2 \times p)$ from the second population. Each row represents a

*p*-dimensional observation.

14 SF2006.GLHT.NABT

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1},\ldots,\boldsymbol{y}_{in_i}$$
, are i. i. d. with  $\mathrm{E}(\boldsymbol{y}_{i1})=\boldsymbol{\mu}_i$ ,  $\mathrm{Cov}(\boldsymbol{y}_{i1})=\boldsymbol{\Sigma}, i=1,2.$ 

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Srivastava and Du (2008) proposed the following test statistic:

$$T_{SD} = \frac{n^{-1}n_1n_2(\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2)^{\top}\boldsymbol{D}_S^{-1}(\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2) - \frac{(n-2)p}{n-4}}{\sqrt{2\left[\operatorname{tr}(\boldsymbol{R}^2) - \frac{p^2}{n-2}\right]c_{p,n}}},$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors,  $D_S$  is the diagonal matrix of sample variance, R is the sample correlation matrix and  $c_{p,n}$  is the adjustment coefficient proposed by Srivastava and Du (2008). They showed that under the null hypothesis,  $T_{SD}$  is asymptotically normally distributed.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Srivastava MS, Du M (2008). "A test for the mean vector with fewer observations than the dimension." *Journal of Multivariate Analysis*, **99**(3), 386–402. doi:10.1016/j.jmva.2006.11.002.

## **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
SD2008.TS.NABT(group1,group2)</pre>
```

SF2006.GLHT.NABT

Normal-approximation-based test for GLHT problem proposed by Srivastava and Fujikoshi (2006)

#### **Description**

Srivastava and Fujikoshi (2006)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data with assuming that underlying covariance matrices are the same.

SF2006.GLHT.NABT 15

#### Usage

SF2006.GLHT.NABT(Y,X,C,n,p)

#### **Arguments**

Y	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
Χ	A known $n \times k$ full-rank design matrix with $\operatorname{rank}(\boldsymbol{X}) = k < n$ .
С	A known matrix of size $q \times k$ with rank $(C) = q < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

#### **Details**

A high-dimensional linear regression model can be expressed as

$$Y = X\Theta + \epsilon$$
.

where  $\Theta$  is a  $k \times p$  unknown parameter matrix and  $\epsilon$  is an  $n \times p$  error matrix.

It is of interest to test the following GLHT problem

$$H_0: C\Theta = \mathbf{0}, \quad \text{vs.} \quad H_1: C\Theta \neq \mathbf{0}.$$

Srivastava and Fujikoshi (2006) proposed the following test statistic:

$$T_{SF} = \left[2q\hat{a}_2(1+(n-k)^{-1}q)\right]^{-1/2} \left[\frac{\text{tr}(\boldsymbol{B})}{\sqrt{p}} - \frac{q}{\sqrt{n-k}} \frac{\text{tr}(\boldsymbol{W})}{\sqrt{(n-k)p}}\right].$$

where  $\boldsymbol{W}$  and  $\boldsymbol{B}$  are the matrix of sum of squares and products due to error and the error, respectively, and  $\hat{a}_2 = [\operatorname{tr}(\boldsymbol{W}^2) - \operatorname{tr}^2(\boldsymbol{W})/(n-k)]/[(n-k-1)(n-k+2)p]$ . They showed that under the null hypothesis,  $T_{SF}$  is asymptotically normally distributed.

## Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Srivastava MS, Fujikoshi Y (2006). "Multivariate analysis of variance with fewer observations than the dimension." *Journal of Multivariate Analysis*, **97**(9), 1927–1940. doi:10.1016/j.jmva.2005.08.010.

16 SKK2013.TSBF.NABT

### **Examples**

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
\verb|group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group|
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
X \leftarrow matrix(c(rep(1,n[1]),rep(0,sum(n)),rep(1,n[2]), rep(0,sum(n)),
             rep(1,n[3]),rep(0,sum(n)),rep(1,n[4])),ncol=k,nrow=sum(n))
q <- k-1
C <- cbind(diag(q),-rep(1,q))</pre>
SF2006.GLHT.NABT(Y,X,C,n,p)
```

SKK2013.TSBF.NABT

Normal-approximation-based test for two-sample BF problem proposed by Srivastava et al. (2013)

## Description

Srivastava et al. (2013)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

### Usage

```
SKK2013.TSBF.NABT(y1, y2)
```

## **Arguments**

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a p-
	dimensional observation.

y2 The data matrix  $(n_2 \times p \text{ from the second population.}$  Each row represents a p-dimensional observation.

YS2012.GLHT.NABT 17

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Srivastava et al. (2013) proposed the following test statistic:

$$T_{SKK} = \frac{(\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2)^{\top} \hat{\boldsymbol{D}}^{-1} (\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2) - p}{\sqrt{2\widehat{\operatorname{Var}}(\hat{q}_n)c_{p,n}}},$$

where  $\bar{y}_i, i=1,2$  are the sample mean vectors,  $\hat{D}=\hat{D}_1/n_1+\hat{D}_2/n_2$  with  $\hat{D}_i, i=1,2$  being the diagonal matrices consisting of only the diagonal elements of the sample covariance matrices.  $\widehat{\text{Var}}(\hat{q}_n)$  is given by equation (1.18) in Srivastava et al. (2013), and  $c_{p,n}$  is the adjustment coefficient proposed by Srivastava et al. (2013). They showed that under the null hypothesis,  $T_{SKK}$  is asymptotically normally distributed.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

### References

Srivastava MS, Katayama S, Kano Y (2013). "A two sample test in high dimensional data." *Journal of Multivariate Analysis*, **114**, 349–358. doi:10.1016/j.jmva.2012.08.014.

#### **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
SKK2013.TSBF.NABT(group1,group2)</pre>
```

YS2012.GLHT.NABT

Normal-approximation-based test for GLHT problem proposed by Yamada and Srivastava (2012)

#### **Description**

Yamada and Srivastava (2012)'test for general linear hypothesis testing (GLHT) problem for highdimensional data with assuming that underlying covariance matrices are the same. 18 YS2012.GLHT.NABT

#### Usage

YS2012.GLHT.NABT(Y,X,C,n,p)

### **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
Χ	A known $n \times k$ full-rank design matrix with $\operatorname{rank}(\boldsymbol{X}) = k < n$ .
С	A known matrix of size $q \times k$ with rank $(C) = q < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i,n_i.$
р	The dimension of data.

#### **Details**

A high-dimensional linear regression model can be expressed as

$$Y = X\Theta + \epsilon$$
.

where  $\Theta$  is a  $k \times p$  unknown parameter matrix and  $\epsilon$  is an  $n \times p$  error matrix.

It is of interest to test the following GLHT problem

$$H_0: C\Theta = \mathbf{0}, \quad \text{vs. } H_1: C\Theta \neq \mathbf{0}.$$

Yamada and Srivastava (2012) proposed the following test statistic:

$$T_{YS} = \frac{(n-k)\operatorname{tr}(S_h \boldsymbol{D}_{S_e}^{-1}) - (n-k)pq/(n-k-2)}{\sqrt{2q[\operatorname{tr}(\boldsymbol{R}^2) - p^2/(n-k)]c_{p,n}}},$$

where  $S_h$  and  $S_e$  are the variation matrices due to the hypothesis and error, respectively, and  $D_{S_e}$  and R are diagonal matrix with the diagonal elements of  $S_e$  and the sample correlation matrix, respectively.  $c_{p,n}$  is the adjustment coefficient proposed by Yamada and Srivastava (2012). They showed that under the null hypothesis,  $T_{YS}$  is asymptotically normally distributed.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

## References

Yamada T, Srivastava MS (2012). "A test for multivariate analysis of variance in high dimension." *Communications in Statistics-Theory and Methods*, **41**(13-14), 2602–2615. doi:10.1080/03610926.2011.581786.

ZGZ2017.GLHT.2cNRT 19

#### **Examples**

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
X \leftarrow matrix(c(rep(1,n[1]),rep(0,sum(n)),rep(1,n[2]), rep(0,sum(n)),rep(1,n[3]),
             rep(0, sum(n)), rep(1, n[4])), ncol=k, nrow=sum(n))
q < - k-1
C <- cbind(diag(q), -rep(1,q))</pre>
YS2012.GLHT.NABT(Y,X,C,n,p)
```

ZGZ2017.GLHT.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for GLHT problem proposed Zhang et al. (2017)

## **Description**

Zhang et al. (2017)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data with assuming that underlying covariance matrices are the same.

#### Usage

```
ZGZ2017.GLHT.2cNRT(Y,G,n,p)
```

#### **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
G	A known full-rank coefficient matrix $(q \times k)$ with rank $(G) < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i,n_i$ .
p	The dimension of data.

#### **Details**

Suppose we have the following k independent high-dimensional samples:

$$\mathbf{y}_{i1}, \dots, \mathbf{y}_{in_i}$$
, are i. i. d. with  $E(\mathbf{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\mathbf{y}_{i1}) = \boldsymbol{\Sigma}$ ,  $i = 1, \dots, k$ .

It is of interest to test the following GLHT problem:

$$H_0: GM = 0$$
, vs.  $H_1: GM \neq 0$ ,

where  $M = (\mu_1, \dots, \mu_k)^{\top}$  is a  $k \times p$  matrix collecting k mean vectors and  $G : q \times k$  is a known full-rank coefficient matrix with rank(G) < k.

Zhang et al. (2017) proposed the following test statistic:

$$T_{ZGZ} = \|\boldsymbol{C}\hat{\boldsymbol{\mu}}\|^2,$$

where  $C = [(\boldsymbol{G}\boldsymbol{D}\boldsymbol{G}^{\top})^{-1/2}\boldsymbol{G}] \otimes \boldsymbol{I}_p$ , and  $\hat{\boldsymbol{\mu}} = (\bar{\boldsymbol{y}}_1^{\top}, \dots, \bar{\boldsymbol{y}}_k^{\top})^{\top}$ , with  $\bar{\boldsymbol{y}}_i, i = 1, \dots, k$  being the sample mean vectors and  $\boldsymbol{D} = \mathrm{diag}(1/n_1, \dots, 1/n_k)$ .

They showed that under the null hypothesis,  $T_{ZGZ}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Guo J, Zhou B (2017). "Linear hypothesis testing in high-dimensional one-way MANOVA." *Journal of Multivariate Analysis*, **155**, 200–216. doi:10.1016/j.jmva.2017.01.002.

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
G \leftarrow cbind(diag(k-1), rep(-1, k-1))
ZGZ2017.GLHT.2cNRT(Y,G,n,p)
```

ZGZ2017.GLHTBF.NABT Normal-approximation-based test for GLHT problem under heteroscedasticity proposed by Zhou et al. (2017)

#### **Description**

Zhou et al. (2017)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data under heteroscedasticity.

## Usage

ZGZ2017.GLHTBF.NABT(Y,G,n,p)

#### **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
G	A known full-rank coefficient matrix $(q \times k)$ with rank $(G) < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i,n_i.$
p	The dimension of data.

#### **Details**

Suppose we have the following k independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, \dots, k$ .

It is of interest to test the following GLHT problem:

$$H_0: GM = 0$$
, vs.  $H_1: GM \neq 0$ ,

where  $M = (\mu_1, \dots, \mu_k)^{\top}$  is a  $k \times p$  matrix collecting k mean vectors and  $G : q \times k$  is a known full-rank coefficient matrix with  $\operatorname{rank}(G) < k$ .

Let  $\bar{y}_i$ ,  $i=1,\ldots,k$  be the sample mean vectors and  $\hat{\Sigma}_i$ ,  $i=1,\ldots,k$  be the sample covariance matrices.

Zhou et al. (2017) proposed the following U-statistic based test statistic:

$$T_{ZGZ} = \|\boldsymbol{C}\hat{\boldsymbol{\mu}}\|^2 - \sum_{i=1}^k h_{ii}\operatorname{tr}(\hat{\boldsymbol{\Sigma}}_i)/n_i,$$

where  $C = [(GDG^{\top})^{-1/2}G] \otimes I_p$ ,  $D = \text{diag}(1/n_1, \dots, 1/n_k)$ , and  $h_{ij}$  is the (i, j)th entry of the  $k \times k$  matrix  $H = G^{\top}(GDG^{\top})^{-1}G$ .

They showed that under the null hypothesis,  $T_{ZGZ}$  is asymptotically normally distributed.

22 ZGZC2020.TS.2cNRT

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhou B, Guo J, Zhang J (2017). "High-dimensional general linear hypothesis testing under heteroscedasticity." *Journal of Statistical Planning and Inference*, **188**, 36–54. doi:10.1016/j.jspi.2017.03.005.

#### **Examples**

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 \leftarrow as.matrix(corneal[44:57, ]) ## unilateral suspect group
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
G \leftarrow cbind(diag(k-1), rep(-1, k-1))
ZGZ2017.GLHTBF.NABT(Y,G,n,p)
```

ZGZC2020.TS.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for two-sample problem proposed by Zhang et al. (2020)

### **Description**

Zhang et al. (2020)'s test for testing equality of two-sample high-dimensional mean vectors with assuming that two covariance matrices are the same.

#### Usage

```
ZGZC2020.TS.2cNRT(y1, y2)
```

## Arguments

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a $p$ -
	dimensional observation.
y2	The data matrix $(n_2 \times p)$ from the second population. Each row represents a

*p*-dimensional observation.

ZWZ2023.TSBF.2cNRT 23

#### **Details**

Suppose we have two independent high-dimensional samples:

$$y_{i1}, \dots, y_{in_i}$$
, are i. i. d. with  $E(y_{i1}) = \mu_i$ ,  $Cov(y_{i1}) = \Sigma$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \boldsymbol{\mu}_1 = \boldsymbol{\mu}_2 \text{ versus } H_1: \boldsymbol{\mu}_1 \neq \boldsymbol{\mu}_2.$$

Zhang et al.(2020) proposed the following test statistic:

$$T_{ZGZC} = \frac{n_1 n_2}{n} \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2,$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors. They showed that under the null hypothesis,  $T_{ZGZC}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

## Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Guo J, Zhou B, Cheng M (2020). "A simple two-sample test in high dimensions based on L 2-norm." *Journal of the American Statistical Association*, **115**(530), 1011–1027. doi:10.1080/01621459.2019.1604366.

## **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZGZC2020.TS.2cNRT(group1, group2)</pre>
```

ZWZ2023.TSBF.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for two-sample BF problem proposed by Zhu et al. (2023)

### Description

Zhu et al. (2023)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

### Usage

```
ZWZ2023.TSBF.2cNRT(y1, y2)
```

#### **Arguments**

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a $p$ -dimensional observation.
y2	The data matrix $(n_2 \times p)$ from the second population. Each row represents a $p$ -dimensional observation.

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\mathbf{y}_{i1}, \dots, \mathbf{y}_{in_i}$$
, are i. i. d. with  $\mathbf{E}(\mathbf{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $\mathbf{Cov}(\mathbf{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Zhu et al. (2023) proposed the following test statistic:

$$T_{ZWZ} = \frac{n_1 n_2 n^{-1} \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2}{\operatorname{tr}(\hat{\boldsymbol{\Omega}}_n)},$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors and  $\hat{\Omega}_n$  is the estimator of  $\text{Cov}[(n_1n_2/n)^{1/2}(\bar{y}_1-\bar{y}_2)]$ . They showed that under the null hypothesis,  $T_{ZWZ}$  and an F-type mixture have the same normal or non-normal limiting distribution.

## Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhu T, Wang P, Zhang J (2023). "Two-sample Behrens–Fisher problems for high-dimensional data: a normal reference F-type test." *Computational Statistics*, 1–24. doi:10.1007/s00180023014336.

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZWZ2023.TSBF.2cNRT(group1, group2)</pre>
```

ZZ2022.GLHT.3cNRT 25

ZZ2022.GLHT.3cNRT	Normal-reference-test with three-cumulant (3-c) matched \$\chi^2\$-
	approximation for GLHT problem proposed by Zhu and Zhang (2022)

### Description

Zhu and Zhang (2022)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data with assuming that underlying covariance matrices are the same.

## Usage

ZZ2022.GLHT.3cNRT(Y,G,n,p)

## **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
G	A known full-rank coefficient matrix $(q \times k)$ with rank $(G) < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

#### **Details**

Suppose we have the following k independent high-dimensional samples:

$$y_{i1}, \ldots, y_{in_i}$$
, are i. i. d. with  $E(y_{i1}) = \mu_i$ ,  $Cov(y_{i1}) = \Sigma$ ,  $i = 1, \ldots, k$ .

It is of interest to test the following GLHT problem:

$$H_0: GM = 0$$
, vs.  $H_1: GM \neq 0$ ,

where  $M = (\mu_1, \dots, \mu_k)^{\top}$  is a  $k \times p$  matrix collecting k mean vectors and  $G : q \times k$  is a known full-rank coefficient matrix with  $\operatorname{rank}(G) < k$ .

Zhu and Zhang (2022) proposed the following test statistic:

$$T_{ZZ} = \|\boldsymbol{C}\hat{\boldsymbol{\mu}}\|^2 - q\operatorname{tr}(\hat{\boldsymbol{\Sigma}}),$$

where  $C = [(\boldsymbol{G}\boldsymbol{D}\boldsymbol{G}^{\top})^{-1/2}\boldsymbol{G}] \otimes \boldsymbol{I}_p$ , and  $\hat{\boldsymbol{\mu}} = (\bar{\boldsymbol{y}}_1^{\top}, \dots, \bar{\boldsymbol{y}}_k^{\top})^{\top}$ , with  $\bar{\boldsymbol{y}}_i, i = 1, \dots, k$  being the sample mean vectors and  $\hat{\boldsymbol{\Sigma}}$  being the usual pooled sample covariance matrix of the k samples.

They showed that under the null hypothesis,  $T_{ZZ}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhu T, Zhang J (2022). "Linear hypothesis testing in high-dimensional one-way MANOVA: a new normal reference approach." *Computational Statistics*, **37**(1), 1–27. doi:10.1007/s0018002101110-6.

## **Examples**

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
G \leftarrow cbind(diag(k-1), rep(-1, k-1))
ZZ2022.GLHT.3cNRT(Y,G,n,p)
```

ZZ2022.GLHTBF.3cNRT

Normal-reference-test with three-cumulant (3-c) matched \$\chi^2\$-approximation for GLHT problem under heteroscedasticity proposed by Zhang and Zhu (2022)

## **Description**

Zhang and Zhu (2022)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data under heteroscedasticity.

### Usage

```
ZZ2022.GLHTBF.3cNRT(Y,G,n,p)
```

#### **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
G	A known full-rank coefficient matrix $(q \times k)$ with rank $(G) < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

ZZ2022.GLHTBF.3cNRT 27

#### **Details**

Suppose we have the following k independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, \dots, k$ .

It is of interest to test the following GLHT problem:

$$H_0: GM = 0$$
, vs.  $H_1: GM \neq 0$ ,

where  $M = (\mu_1, \dots, \mu_k)^{\top}$  is a  $k \times p$  matrix collecting k mean vectors and  $G : q \times k$  is a known full-rank coefficient matrix with rank(G) < k.

Let  $\bar{y}_i$ ,  $i=1,\ldots,k$  be the sample mean vectors and  $\hat{\Sigma}_i$ ,  $i=1,\ldots,k$  be the sample covariance matrices.

Zhang and Zhu (2022) proposed the following U-statistic based test statistic:

$$T_{ZZ} = \|\boldsymbol{C}\hat{\boldsymbol{\mu}}\|^2 - \sum_{i=1}^k h_{ii} \operatorname{tr}(\hat{\boldsymbol{\Sigma}}_i)/n_i,$$

where  $C = [(GDG^{\top})^{-1/2}G] \otimes I_p$ ,  $D = \text{diag}(1/n_1, \dots, 1/n_k)$ , and  $h_{ij}$  is the (i, j)th entry of the  $k \times k$  matrix  $H = G^{\top}(GDG^{\top})^{-1}G$ .

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Zhu T (2022). "A new normal reference test for linear hypothesis testing in high-dimensional heteroscedastic one-way MANOVA." *Computational Statistics & Data Analysis*, **168**, 107385. doi:10.1016/j.csda.2021.107385.

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
G \leftarrow cbind(diag(k-1), rep(-1, k-1))
ZZ2022.GLHTBF.3cNRT(Y,G,n,p)
```

28 ZZ2022.TS.3cNRT

ZZ2022.TS.3cNRT	Normal-reference-test with three-cumulant (3-c) matched \$\chi^2\$-
	approximation for two-sample problem proposed by Zhang and Zhu (2022)

#### **Description**

Zhang and Zhu (2022)'s test for testing equality of two-sample high-dimensional mean vectors with assuming that two covariance matrices are the same.

#### Usage

```
ZZ2022.TS.3cNRT(y1, y2)
```

## **Arguments**

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a $p$ -
	dimensional observation

y2 The data matrix  $(n_2 \times p)$  from the second population. Each row represents a p-dimensional observation.

#### **Details**

Suppose we have two independent high-dimensional samples:

$$y_{i1}, \dots, y_{in_i}$$
, are i. i. d. with  $E(y_{i1}) = \mu_i$ ,  $Cov(y_{i1}) = \Sigma$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Zhang et al.(2022) proposed the following test statistic:

$$T_{ZZ} = \frac{n_1 n_2}{n} \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2 - \operatorname{tr}(\hat{\boldsymbol{\Sigma}}),$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors and  $\hat{\Sigma}$  is the pooled sample covariance matrix. They showed that under the null hypothesis,  $T_{ZZ}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Zhu T (2022). "A revisit to Bai–Saranadasa's two-sample test." *Journal of Nonparametric Statistics*, **34**(1), 58–76. doi:10.1080/10485252.2021.2015768.

ZZ2022.TSBF.3cNRT 29

#### **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZZ2022.TS.3cNRT(group1, group2)</pre>
```

ZZ2022.TSBF.3cNRT

Normal-reference-test with three-cumulant (3-c) matched \$\chi^2\$-approximation for two-sample BF problem proposed by Zhang and Zhu (2022)

#### **Description**

Zhang and Zhu (2022)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

## Usage

```
ZZ2022.TSBF.3cNRT(y1, y2)
```

### **Arguments**

y1 The data matrix  $(n_1 \times p)$  from the first population. Each row represents a p-dimensional observation.

y2 The data matrix  $(n_2 \times p)$  from the second population. Each row represents a p-dimensional observation.

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1}, \dots, \boldsymbol{y}_{in_i}$$
, are i. i. d. with  $E(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i$ ,  $Cov(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i$ ,  $i = 1, 2$ .

The primary object is to test

$$H_0: \boldsymbol{\mu}_1 = \boldsymbol{\mu}_2 \text{ versus } H_1: \boldsymbol{\mu}_1 \neq \boldsymbol{\mu}_2.$$

Zhang and Zhu (2022) proposed the following test statistic:

$$T_{ZZ} = \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2 - \operatorname{tr}(\hat{\boldsymbol{\Omega}}_n),$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors and  $\hat{\Omega}_n$  is the estimator of  $\text{Cov}(\bar{y}_1 - \bar{y}_2)$ . They showed that under the null hypothesis,  $T_{ZZ}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

## Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Zhu T (2022). "A further study on Chen-Qin's test for two-sample Behrens–Fisher problems for high-dimensional data." *Journal of Statistical Theory and Practice*, **16**(1), 1. doi:10.1007/s4251902100232w.

## **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZZ2022.TSBF.3cNRT(group1, group2)</pre>
```

ZZG2022.GLHTBF.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for GLHT problem under heteroscedasticity proposed by Zhang et al. (2022)

## **Description**

Zhang et al. (2022)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data under heteroscedasticity.

## Usage

```
ZZG2022.GLHTBF.2cNRT(Y,G,n,p)
```

## Arguments

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
G	A known full-rank coefficient matrix $(q \times k)$ with rank $(G) < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

#### **Details**

Suppose we have the following k independent high-dimensional samples:

$$y_{i1}, \ldots, y_{in_i}$$
, are i. i. d. with  $E(y_{i1}) = \mu_i$ ,  $Cov(y_{i1}) = \Sigma_i$ ,  $i = 1, \ldots, k$ .

It is of interest to test the following GLHT problem:

$$H_0: GM = 0$$
, vs.  $H_1: GM \neq 0$ ,

where  $M = (\mu_1, \dots, \mu_k)^{\top}$  is a  $k \times p$  matrix collecting k mean vectors and  $G : q \times k$  is a known full-rank coefficient matrix with  $\operatorname{rank}(G) < k$ .

Zhang et al. (2022) proposed the following test statistic:

$$T_{ZZG} = \|\boldsymbol{C}\hat{\boldsymbol{\mu}}\|^2,$$

where 
$$C = [(\boldsymbol{G}\boldsymbol{D}\boldsymbol{G}^{\top})^{-1/2}\boldsymbol{G}] \otimes \boldsymbol{I}_p$$
 with  $\boldsymbol{D} = \operatorname{diag}(1/n_1,\ldots,1/n_k)$ , and  $\hat{\boldsymbol{\mu}} = (\bar{\boldsymbol{y}}_1^{\top},\ldots,\bar{\boldsymbol{y}}_k^{\top})^{\top}$  with  $\bar{\boldsymbol{y}}_i, i=1,\ldots,k$  being the sample mean vectors.

They showed that under the null hypothesis,  $T_{ZZG}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Zhou B, Guo J (2022). "Linear hypothesis testing in high-dimensional heteroscedastic one-way MANOVA: A normal reference  $L^2$ -norm based test." *Journal of Multivariate Analysis*, **187**, 104816. doi:10.1016/j.jmva.2021.104816.

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
G \leftarrow cbind(diag(k-1), rep(-1, k-1))
ZZG2022.GLHTBF.2cNRT(Y,G,n,p)
```

ZZGZ2021.TSBF.2cNRT

ZZGZ2021.TSBF.2cNRT Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for two-sample BF problem proposed by Zhang et al. (2021)

#### **Description**

Zhang et al. (2021)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

### Usage

ZZGZ2021.TSBF.2cNRT(y1, y2)

#### **Arguments**

y1 The data matrix  $(n_1 \times p)$  from the first population. Each row represents a p-dimensional observation.

y2 The data matrix  $(n_2 \times p)$  from the second population. Each row represents a p-dimensional observation.

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1},\ldots,\boldsymbol{y}_{in_i}$$
, are i. i. d. with  $\mathrm{E}(\boldsymbol{y}_{i1})=\boldsymbol{\mu}_i$ ,  $\mathrm{Cov}(\boldsymbol{y}_{i1})=\boldsymbol{\Sigma}_i, i=1,2.$ 

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Zhang et al.(2021) proposed the following test statistic:

$$T_{ZZGZ} = \frac{n_1 n_2}{n} \|\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2\|^2,$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors. They showed that under the null hypothesis,  $T_{ZZGZ}$  and a chi-squared-type mixture have the same normal or non-normal limiting distribution.

### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang J, Zhou B, Guo J, Zhu T (2021). "Two-sample Behrens-Fisher problems for high-dimensional data: A normal reference approach." *Journal of Statistical Planning and Inference*, **213**, 142–161. doi:10.1016/j.jspi.2020.11.008.

ZZZ2020.TS.2cNRT 33

#### **Examples**

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZZGZ2021.TSBF.2cNRT(group1, group2)</pre>
```

ZZZ2020.TS.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for two-sample problem proposed by Zhang et al. (2020)

## Description

Zhang et al. (2020)'s test for testing equality of two-sample high-dimensional mean vectors with assuming that two covariance matrices are the same.

#### Usage

```
ZZZ2020.TS.2cNRT(y1, y2)
```

### **Arguments**

y1 The data matrix  $(n_1 \times p)$  from the first population. Each row represents a p-dimensional observation.

y2 The data matrix  $(n_2 \times p)$  from the second population. Each row represents a p-dimensional observation.

#### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1},\ldots,\boldsymbol{y}_{in_i}$$
, are i. i. d. with  $\mathrm{E}(\boldsymbol{y}_{i1})=\boldsymbol{\mu}_i$ ,  $\mathrm{Cov}(\boldsymbol{y}_{i1})=\boldsymbol{\Sigma}, i=1,2.$ 

The primary object is to test

$$H_0: \boldsymbol{\mu}_1 = \boldsymbol{\mu}_2 \text{ versus } H_1: \boldsymbol{\mu}_1 \neq \boldsymbol{\mu}_2.$$

Zhang et al.(2020) proposed the following test statistic:

$$T_{ZZZ} = \frac{n_1 n_2}{np} (\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2)^\top \hat{\boldsymbol{D}}^{-1} (\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2),$$

where  $\bar{y}_i$ , i=1,2 are the sample mean vectors,  $\hat{D}$  is the diagonal matrix of sample covariance matrix. They showed that under the null hypothesis,  $T_{ZZZ}$  and a chi-squared-type mixture have the same limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhang L, Zhu T, Zhang J (2020). "A simple scale-invariant two-sample test for high-dimensional data." *Econometrics and Statistics*, **14**, 131–144. doi:10.1016/j.ecosta.2019.12.002.

## Examples

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZZZ2020.TS.2cNRT(group1,group2)</pre>
```

ZZZ2022.GLHT.2cNRT

Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-approximation for GLHT problem proposed by Zhu et al. (2022)

## Description

Zhu et al. (2022)'s test for general linear hypothesis testing (GLHT) problem for high-dimensional data with assuming that underlying covariance matrices are the same.

## Usage

```
ZZZ2022.GLHT.2cNRT(Y,X,C,n,p)
```

## **Arguments**

Υ	A list of $k$ data matrices. The $i$ th element represents the data matrix $(n_i \times p)$ from the $i$ th population with each row representing a $p$ -dimensional observation.
Χ	A known $n \times k$ full-rank design matrix with $\operatorname{rank}(\boldsymbol{X}) = k < n$ .
С	A known matrix of size $q \times k$ with $rank(\mathbf{C}) = q < k$ .
n	A vector of $k$ sample sizes. The $i$ th element represents the sample size of group $i, n_i$ .
р	The dimension of data.

ZZZ2022.GLHT.2cNRT 35

#### **Details**

A high-dimensional linear regression model can be expressed as

$$Y = X\Theta + \epsilon$$
,

where  $\Theta$  is a  $k \times p$  unknown parameter matrix and  $\epsilon$  is an  $n \times p$  error matrix.

It is of interest to test the following GLHT problem

$$H_0: C\Theta = \mathbf{0}$$
, vs.  $H_1: C\Theta \neq \mathbf{0}$ .

Zhu et al. (2022) proposed the following test statistic:

$$T_{ZZZ} = \frac{(n-k-2)}{(n-k)pq} \operatorname{tr}(\boldsymbol{S}_h \boldsymbol{D}^{-1}),$$

where  $S_h$  and  $S_e$  are the variation matrices due to the hypothesis and error, respectively, and D is the diagonal matrix with the diagonal elements of  $S_e/(n-k)$ . They showed that under the null hypothesis,  $T_{ZZZ}$  and a chi-squared-type mixture have the same limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

#### References

Zhu T, Zhang L, Zhang J (2023). "Hypothesis Testing in High-Dimensional Linear Regression: A Normal Reference Scale-Invariant Test." *Statistica Sinica*. doi:10.5705/ss.202020.0362.

```
library("HDNRA")
data("corneal")
dim(corneal)
group1 <- as.matrix(corneal[1:43, ]) ## normal group</pre>
group2 <- as.matrix(corneal[44:57, ]) ## unilateral suspect group</pre>
group3 <- as.matrix(corneal[58:78, ]) ## suspect map group</pre>
group4 <- as.matrix(corneal[79:150, ]) ## clinical keratoconus group</pre>
p <- dim(corneal)[2]</pre>
Y <- list()
k <- 4
Y[[1]] <- group1
Y[[2]] <- group2
Y[[3]] <- group3
Y[[4]] <- group4
n <- c(nrow(Y[[1]]),nrow(Y[[2]]),nrow(Y[[3]]),nrow(Y[[4]]))</pre>
X \leftarrow matrix(c(rep(1,n[1]),rep(0,sum(n)),rep(1,n[2]), rep(0,sum(n)),
             rep(1,n[3]),rep(0,sum(n)),rep(1,n[4])),ncol=k,nrow=sum(n))
q <- k-1
C <- cbind(diag(q),-rep(1,q))</pre>
ZZZ2022.GLHT.2cNRT(Y,X,C,n,p)
```

36 ZZZ2023.TSBF.2cNRT

ZZZ2023.TSBF.2cNRT	Normal-reference-test with two-cumulant (2-c) matched \$\chi^2\$-
	approximation for two-sample BF problem proposed by Zhang et al. (2023)

## **Description**

Zhang et al. (2023)'s test for testing equality of two-sample high-dimensional mean vectors without assuming that two covariance matrices are the same.

#### Usage

```
ZZZ2023.TSBF.2cNRT(y1, y2, cutoff)
```

## Arguments

y1	The data matrix $(n_1 \times p)$ from the first population. Each row represents a $p$ -dimensional observation.
y2	The data matrix $(n_2 \times p)$ from the second population. Each row represents a $p$ -dimensional observation.
cutoff	An empirical criterion for applying the adjustment coefficient

### **Details**

Suppose we have two independent high-dimensional samples:

$$\boldsymbol{y}_{i1},\ldots,\boldsymbol{y}_{in_i}, \text{ are i. i. d. with } \mathrm{E}(\boldsymbol{y}_{i1}) = \boldsymbol{\mu}_i, \ \mathrm{Cov}(\boldsymbol{y}_{i1}) = \boldsymbol{\Sigma}_i, i = 1, 2.$$

The primary object is to test

$$H_0: \mu_1 = \mu_2 \text{ versus } H_1: \mu_1 \neq \mu_2.$$

Zhang et al.(2023) proposed the following test statistic:

$$T_{ZZZ} = \frac{n_1 n_2}{np} (\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2)^\top \hat{\boldsymbol{D}}_n^{-1} (\bar{\boldsymbol{y}}_1 - \bar{\boldsymbol{y}}_2),$$

where  $\bar{\boldsymbol{y}}_i, i=1,2$  are the sample mean vectors, and  $\hat{\boldsymbol{D}}_n=\mathrm{diag}(\hat{\boldsymbol{\Sigma}}_1/n+\hat{\boldsymbol{\Sigma}}_2/n)$  with  $n=n_1+n_2$ . They showed that under the null hypothesis,  $T_{ZZZ}$  and a chi-squared-type mixture have the same limiting distribution.

#### Value

A list of class "NRtest" containing the results of the hypothesis test. See the help file for NRtest.object for details.

ZZZ2023.TSBF.2cNRT 37

## References

Zhang L, Zhu T, Zhang J (2023). "Two-sample Behrens–Fisher problems for high-dimensional data: a normal reference scale-invariant test." *Journal of Applied Statistics*, **50**(3), 456–476. doi:10.1080/02664763.2020.1834516.

```
library("HDNRA")
data("COVID19")
dim(COVID19)
group1 <- as.matrix(COVID19[c(2:19, 82:87), ]) ## healthy group
group2 <- as.matrix(COVID19[-c(1:19, 82:87), ]) ## COVID-19 patients
ZZZ2023.TSBF.2cNRT(group1,group2,cutoff=1.2)</pre>
```

# **Index**

```
* datasets
                                                 FHW2004.GLHT.NABT, 7
    corneal, 4
                                                 NRtest.object, 4, 6, 8, 9, 12–15, 17, 18, 20,
    COVID19, 5
                                                          22-25, 27, 28, 30-32, 34-36
* data
    corneal, 4
                                                 print.NRtest, 10, 11
    COVID19, 5
* glht
                                                 S2007.ks.NABT, 12
    FHW2004.GLHT.NABT, 7
                                                 SD2008.TS.NABT, 13
    S2007.ks.NABT, 12
                                                 SF2006.GLHT.NABT, 14
    SF2006.GLHT.NABT, 14
                                                 SKK2013.TSBF.NABT, 16
    YS2012.GLHT.NABT, 17
    ZGZ2017.GLHTBF.NABT, 21
                                                 YS2012.GLHT.NABT, 17
* nraglht
    ZGZ2017.GLHT.2cNRT, 19
                                                 ZGZ2017.GLHT.2cNRT, 19
    ZZ2022.GLHT.3cNRT, 25
                                                 ZGZ2017.GLHTBF.NABT, 21
    ZZ2022.GLHTBF.3cNRT, 26
                                                 ZGZC2020.TS.2cNRT, 22
    ZZG2022.GLHTBF.2cNRT, 30
                                                 ZWZ2023.TSBF.2cNRT, 23
    ZZZ2022.GLHT.2cNRT, 34
                                                 ZZ2022.GLHT.3cNRT, 25
* nrats
                                                 ZZ2022.GLHTBF.3cNRT, 26
    ZGZC2020.TS.2cNRT, 22
                                                 ZZ2022.TS.3cNRT, 28
    ZWZ2023.TSBF.2cNRT, 23
                                                 ZZ2022.TSBF.3cNRT, 29
    ZZ2022.TS.3cNRT, 28
                                                 ZZG2022.GLHTBF.2cNRT, 30
    ZZ2022.TSBF.3cNRT, 29
                                                 ZZGZ2021.TSBF.2cNRT, 32
    ZZGZ2021.TSBF.2cNRT, 32
                                                 ZZZ2020.TS.2cNRT, 33
    ZZZ2020.TS.2cNRT, 33
                                                 ZZZ2022.GLHT.2cNRT, 34
    ZZZ2023.TSBF.2cNRT, 36
                                                 ZZZ2023.TSBF.2cNRT, 36
* object
    NRtest.object, 9
    print.NRtest, 11
* ts
    BS1996.TS.NABT, 3
    CQ2010.TSBF.NABT, 6
    SD2008.TS.NABT, 13
    SKK2013.TSBF.NABT, 16
BS1996.TS.NABT, 3
{\tt corneal}, {\tt 4}
COVID19, 5
CQ2010.TSBF.NABT, 6
```