# Package 'tWLSA'

## January 1, 2021

Title Two way Kernel Weighted Least Squate Approach for Missing Value

Type Package

| Imputation of Metabol  | lomics Data  |  |  |                    |     |
|--|--|--|--|--------------------|-----|
| Version 1.0  |  |  |  |                    |     |
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| Author Nishith Kumar, Md. Aminul Hoque and Masahiro Sugimoto  Maintainer Nishith Kumar <nk.bru09@gmail.com>  Description Missing value imputation technique by minimizing two- way kernel weighted square error loss function</nk.bru09@gmail.com> |  |  |  |                    |     |
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|  |  |  |  | R topics document  | ed: |
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#### **Description**

This package implements a Kernel weight based two way weighted least square approach for missing value imputation of metabolomics data

Package: tWLSA
Type: Package
Version: 1.0
Date: 2021-01-01
License: GPL
Depends: R (>=2.10)

Package tWLSA has the following functions:

chkMiss(): This function check the missing values in the dataset

function.

chkOutliers(): This function Checks row wise outliers in a data matrix

function.

chkOutMiss(): This function Checks both outliers and missing values data matrix

function.

missChkOut(): This function checks the outliers

function.

removeOut(): This function clean the outliers

function.

wlsMisImp(): This function impute the missing values using two way kernel weight based least square approach

Nishith Kumar

Maintainer: Nishith Kumar <nk.bru09@gmail.com>

package

chkMiss

Checking Missing cell in a data matrix

#### **Description**

This function will check whether the data matrix contains missing values or not.

#### Usage

chkMiss(x)

#### **Arguments**

x Metabolomics Data Matrix, whose row contains metabolites and column con-

tains subjects

#### Value

Provide message about the missing values in the data matrix

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#### Author(s)

Nishith Kumar <nk.bru09@gmail.com>

## **Examples**

```
data(dummyDataMiss)
chkMiss(dummyDataMiss)
```

chkOutliers

Checking row wise outliers in a data matrix

## Description

This function check the row-wise outliers in the data matrix using IQR rule.

## Usage

```
chkOutliers(x)
```

## **Arguments**

Х

Metabolomics data matrix

## Details

Checking the row-wise outliers in the data matrix using IQR rule

## Value

Provide the information whether the data matrix contain outliers or not.

#### Author(s)

Nishith Kumar <nk.bru09@gmail.com>

```
data(dummyDataFull)
chkOutliers(dummyDataFull)
```

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chkOutMiss

Checking outliers and missing values data matrix

#### **Description**

Checking outliers and missing values data matrix

#### Usage

```
chkOutMiss(x)
```

#### **Arguments**

Х

Metabolomics data matrix

#### **Details**

This function will Check both the outliers and missing values data matrix

## Value

Provide the information whether the data matrix contain both the outliers and missing values or not.

#### Author(s)

Nishith Kumar <nk.bru09@gmail.com>

## **Examples**

```
data(dummyDataMiss)
chkOutMiss(dummyDataMiss)
```

dummyDataFull

Dummy Metabolomics Data matrix

## Description

Dummy Metabolomics Data matrix with no missing values

## Usage

```
data("dummyDataFull")
```

#### **Format**

```
The format is: num [1:40, 1:45] 0.678 0.671 0.651 5.645 2.437 ... - attr(*, "dimnames")=List of 2 ...$ : chr [1:40] "Guanidoacetate" "O-Acetylcarnitine" "2-Aminobutyrate" "Formate" ... ...$ : chr [1:45] "Cancer_1" "Cancer_2" "Cancer_3" "Cancer_4" ...
```

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#### **Details**

Dummy Metabolomics Data matrix with no missing values

#### **Source**

**Dummy Data** 

## **Examples**

```
data(dummyDataFull)
## maybe str(dummyDataFull) ; plot(dummyDataFull) ...
```

dummyDataMiss

Dummy Metabolomics Dataset with Missing values

## Description

Dummy Metabolomics Dataset with Missing values

## Usage

```
data("dummyDataMiss")
```

#### **Format**

```
The format is: num [1:40, 1:45] 0.678 NA 0.651 5.645 NA ... - attr(*, "dimnames")=List of 2 ..$ : chr [1:40] "Guanidoacetate" "O-Acetylcarnitine" "2-Aminobutyrate" "Formate" ... ..$ : chr [1:45] "Cancer_1" "Cancer_2" "Cancer_3" "Cancer_4" ...
```

## **Details**

Dummy Metabolomics Dataset with Missing values

```
data(dummyDataMiss)
## maybe str(dummyDataMiss) ; plot(dummyDataMiss) ...
```

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missChkOut

Outliers Checking in Dataset

## Description

Checking Outliers using IQR rule

## Usage

```
missChkOut(x)
```

## Arguments

Х

Metabolomics Data Matrix

#### Value

0-indicate no outliers; 1-indicates the existence of outliers

#### Author(s)

Nishith Kumar <nk.bru09@gmail.com>

## **Examples**

```
data(dummyDataFull)
missChkOut(dummyDataFull)
```

removeOut

Data cleaning

## Description

Checking Outliers using IQR rule and replaced it into the group median

#### Usage

```
removeOut(x, group = 1, nGrpMem = ncol(x))
```

## Arguments

x Metabolomics Data Matrix
 group No of group in the Data matrix
 nGrpMem number of sbuject in each group

## Author(s)

Nishith Kumar <nk.bru09@gmail.com>

```
data(dummyDataFull)
modiData<-removeOut(dummyDataFull,group=2,nGrpMem=c(28,17))</pre>
```

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| wlsMisImp | Two way kernel weight based least square approach for missing value imputation |
|-----------|--|
|           |  |

## Description

Two way kernel weight based least square approach for missing value imputation

## Usage

```
wlsMisImp(x, lamda = 0.1, conRate = 99, group = 1, noGrpMem = c(ncol(x)), outlier = TRUE)
```

## Arguments

| X        | Metabolomics Data Matrix  |
|----------|---|
| lamda    | Tuning parameter. Default is 0.1  |
| conRate  | Percent of variation explained by the modiffied dataset. Default is 99. |
| group    | Number of group in the dataset. Default is 1.                           |
| noGrpMem | Number of subject of each group.  |
| outlier  | Outlier condition TRUE/FALSE.   |

## Value

Reconstructed data matrix after inputing missing values

## Author(s)

Nishith Kumar<nk.bru09@gmail.com>

```
data(dummyDataMiss)
recData<-wlsMisImp(dummyDataMiss,lamda=0.1,conRate=99,group=2,noGrpMem=c(28,17),outlier=TRUE)</pre>
```

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