# Package 'opdisDownsampling'

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Type Package	
Title Optimal Distribution Preserving Down-Sampling of Bio-Medical Data	
Version 1.0.1	
<b>Description</b> An optimized method for distribution-preserving class-proportional downsampling of bio-medical data.	
<b>Depends</b> R (>= $3.5.0$ )	
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FlowcytometricData

Example data of hematologic marker expression.

### **Description**

Data set of 6 flow cytometry-based lymphoma makers from 55,843 cells from healthy subjects (class 1) and 55,843 cells from lymphoma patients (class 2).

## Usage

```
data("FlowcytometricData")
```

#### **Details**

Size  $111686 \times 6$ , stored in FlowcytometricData\$[Var\_1,Var\_2,Var\_3,Var\_4,Var\_5,Var\_6] Classes 2, stored in FlowcytometricData\$Cls

#### **Examples**

```
data(FlowcytometricData)
str(FlowcytometricData)
```

GMMartificialData

Example data an artificial Gaussian mixture.

## **Description**

Dataset of 30000 instances with 10 variables that are Gaussian mixtures and belong to classes Cls = 1, 2, or 3, with different means and standard deviations and equal weights of 0.5, 0.4, and 0.1, respectively.

#### Usage

```
data("GMMartificialData")
```

#### **Details**

```
Size 30000 \times 10, stored in GMMartificialData[X1,X2,X3,X4,X5,X6,X7,X8,X9,X10] Classes 3, stored in GMMartificialDataCls
```

## **Examples**

```
data(GMMartificialData)
str(GMMartificialData)
```

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opdisDownsampling Optimal Distribution	on Preserving Down-Sampling of Bio-Medical Data
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## Description

The package provides the necessary functions for optimal distribution-preserving down-sampling of large (bio-medical) data sets.

## Usage

```
opdisDownsampling(Data, Cls, Size, Seed, nTrials = 1000,
TestStat = "ad", MaxCores = getOption("mc.cores", 2L), PCAimportance = FALSE)
```

## Arguments

Data	the (numerical!) data as a vector, matrix or data frame.
Cls	the class information, if any, as a vector of similar length as instances in the data.
Size	the total number of instances across all classes to be drawn.
Seed	a predefined seed to modify the results.
nTrials	how many samples to choose from should be randomly drawn.
TestStat	statistical criterion for similarity judgment.
MaxCores	maximum number of cpu cores to use for parallel computing.
PCAimportance	PCA based feature selection; only variables important in PCA projection are

## Value

Returns a list of data containing the drawn samples and the omitted data.

considered.

ReducedData the selected sample data and class information.

ReducedData the not-selected sample data and class information.

ReducedInstances

the instance numbers of the selected sample data.

#### Author(s)

Jorn Lotsch

### References

Lotsch, J., Malkusch, S., Ultsch, A. (2021): Optimal distribution-preserving downsampling of large biomedical data sets (opdisDownsampling). PLoS One. 2021 Aug 5;16(8):e0255838. doi: 10.1371/journal.pone.0255838. eCollection 2021.

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## Examples

```
## example 1
data(iris)
Iris50percent <- opdisDownsampling(Data = iris[,1:4], Cls = as.integer(iris$Species),
    Size = 50, MaxCores = 1)</pre>
```

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