

Package: a4Core (via r-universe)

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Type Package

Title Automated Affymetrix Array Analysis Core Package

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Description Utility functions for the Automated Affymetrix Array Analysis set of packages.

Imports Biobase, glmnet, methods, stats

Suggests knitr, rmarkdown

License GPL-3

biocViews Microarray, Classification

RoxygenNote 7.1.1

VignetteBuilder knitr

Repository <https://bioc.r-universe.dev>

RemoteUrl <https://github.com/bioc/a4Core>

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confusionMatrix	<i>Generic function to produce a confusion matrix (related to a classification problem)</i>
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Description

Generic function to produce a confusion matrix (related to a classification problem)

Usage

```
confusionMatrix(x, ...)
```

Arguments

x	object (usually a model fit object) that contains all information needed to produce the confusion matrix.
...	further arguments for a specific method

Value

A confusion matrix

Author(s)

Tobias Verbeke

simulateData	<i>Simulate Data for Package Testing and Demonstration Purposes</i>
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Description

Simulate Data for Package Testing and Demonstration Purposes

Usage

```
simulateData(nCols = 40, nRows = 1000, nEffectRows = 5, nNoEffectCols = 5,  
  betweenClassDifference = 1, withinClassSd = 0.5)
```

Arguments

nCols	number of samples; currently this should be an even number
nRows	number of features (genes)
nEffectRows	number of differentially expressed features
nNoEffectCols	number of samples for which the profile of a differentially expressed feature will be set similar to the other class
betweenClassDifference	Average mean difference between the two classes to simulate a certain signal in the features for which an effect was introduced; the default is set to 1
withinClassSd	Within class standard deviation used to add a certain noise level to the features for which an effect was introduced; the default standard deviation is set to 0.5

Value

object of class ExpressionSet with the characteristics specified

Note

The simulation assumes the variances are equal between the two classes. Heterogeneity could easily be introduced in the simulation if this would be requested by the users.

Author(s)

W. Talloen and T. Verbeke

Examples

```
someEset <- simulateData(nCols = 40, nRows = 1000, nEffectRows = 5, nNoEffectCols = 5)
someEset
```

topTable	<i>S4 Generic for obtaining a top table</i>
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Description

a top table is a rectangular object (e.g. data frame) which lists the top n most relevant variables

Usage

```
topTable(fit, n, ...)
```

Arguments

fit	object for which to obtain a top table, generally a fit object for a given model class
n	number of features (variables) to list in the top table, ranked by importance
...	further arguments for specific methods

Value

Top table with top n relevant variable.

Author(s)

Tobias Verbeke

topTable-methods

Methods for topTable

Description

Methods for topTable. topTable extracts the top n most important features for a given classification or regression procedure

Arguments

fit	object resulting from a classification or regression procedure
n	number of features that one wants to extract from a table that ranks all features according to their importance in the classification or regression model; defaults to 10 for limma objects

Methods

glmnet and lognet

fit = "glmnet", n = "numeric" glmnet objects are produced by lassoClass (a4Classif) or lassoReg (a4Base)

- fit = "lognet", n = "numeric" lognet objects are produced by lassoClass (a4Classif) or lassoReg (a4Base)
- fit = "elnet", n = "numeric" elnet objects are produced by lassoClass (a4Classif) or lassoReg (a4Base)

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