

# Package ‘a4Classif’

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**Type** Package

**Title** Automated Affymetrix Array Analysis Classification Package

**Version** 1.56.0

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**Description** Functionalities for classification of Affymetrix microarray data, integrating within the Automated Affymetrix Array Analysis set of packages.

**Depends** a4Core, a4Preproc

**Imports** methods, Biobase, ROCR, pamr, glmnet, varSelRF, utils, graphics, stats

**Suggests** ALL, hgu95av2.db, knitr, rmarkdown

**License** GPL-3

**biocViews** Microarray, GeneExpression, Classification

**VignetteBuilder** knitr

**RoxygenNote** 7.1.1

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<b>lassoClass</b>	<i>Classify using the Lasso</i>
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## Description

Classify using the Lasso

## Usage

```
lassoClass(object, groups)
```

## Arguments

- |        |   |
|--------|---|
| object | object containing the expression measurements; currently the only method supported is one for ExpressionSet objects |
| groups | character string indicating the column containing the class membership  |

## Value

object of class `glmnet`

## Author(s)

Willem Talloen

## References

Goehlmann, H. and W. Talloen (2009). Gene Expression Studies Using Affymetrix Microarrays, Chapman \& Hall/CRC, pp. 183, 205 and 212.

## See Also

[glmnet](#)

## Examples

```
if (require(ALL)){
  data(ALL, package = "ALL")
  ALL <- addGeneInfo(ALL)
  ALL$BTtype <- as.factor(substr(ALL$BT,0,1))

  resultLasso <- lassoClass(object = ALL, groups = "BTtype")
  plot(resultLasso, label = TRUE,
       main = "Lasso coefficients in relation to degree of
       penalization.")
  topTable(resultLasso, n = 15)
}
```

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pamClass

*Classify using Prediction Analysis for MicroArrays*

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## Description

Classify using the Prediction Analysis for MicroArrays (PAM) algorithm as implemented in the pamr package

## Usage

```
pamClass(object, groups, probe2gene = TRUE)
```

## Arguments

object	object containing the expression measurements; currently the only method supported is one for ExpressionSet objects
groups	character string indicating the column containing the class membership
probe2gene	logical; if TRUE Affymetrix probeset IDs are translated into gene symbols; if FALSE no such translation is conducted

## Value

object of class pamClass

## Author(s)

Willem Talloen

## References

Robert Tibshirani, Trevor Hastie, Balasubramanian Narasimhan, and Gilbert Chu (1999). Diagnosis of multiple cancer types by shrunken centroids of gene expression. PNAS 99: 6567-6572. Available at [www.pnas.org](http://www.pnas.org)

Goehlmann, H. and W. Talloen (2009). Gene Expression Studies Using Affymetrix Microarrays, Chapman \& Hall/CRC, p. 221.

**See Also**

[pamr.train](#)

**Examples**

```
if(require(ALL)){
  data(ALL, package = "ALL")
  ALL <- addGeneInfo(ALL)
  ALL$BTtype <- as.factor(substr(ALL$BT, 0, 1))
  resultPam <- pamClass(object = ALL, groups = "BTtype")
  plot(resultPam)
  topTable(resultPam, n = 5)
  confusionMatrix(resultPam)
}
```

**rfClass**

*Classify using Random Forests*

**Description**

Classify using the Random Forest algorithm of Breiman (2001)

**Usage**

```
rfClass(object, groups, probe2gene = TRUE)
```

**Arguments**

<b>object</b>	object containing the expression measurements; currently the only method supported is one for ExpressionSet objects
<b>groups</b>	character string indicating the column containing the class membership
<b>probe2gene</b>	logical; if TRUE Affymetrix probeset IDs are translated into gene symbols in the output object; if FALSE no such translation is conducted

**Value**

Object of class 'rfClass'

**Note**

topTable and plot methods are available for 'rfClass' objects.

**Author(s)**

Tobias Verbeke and Willem Talloen

## References

Breiman, L. (2001), *Random Forests*, Machine Learning 45(1), 5-32.

## See Also

[randomForest](#)

## Examples

```
if(require(ALL)){
  data(ALL, package = "ALL")
  ALL <- addGeneInfo(ALL)
  ALL$BTtype <- as.factor(substr(ALL$BT,0,1))
  # select only a subset of the data for computation time reason
  ALLSubset <- ALL[sample.int(n = nrow(ALL), size = 100, replace = TRUE), ]
  resultRf <- rfClass(object = ALLSubset, groups = "BTtype")
  plot(resultRf)
  topTable(resultRf, n = 15)
}
```

ROCCurve

*Receiver operating curve*

## Description

A ROC curve plots the fraction of true positives (TPR = true positive rate) versus the fraction of false positives (FPR = false positive rate) for a binary classifier when the discrimination threshold is varied. Equivalently, one can also plot sensitivity versus (1 - specificity).

## Usage

```
ROCCurve(
  object,
  groups,
  probesetId = NULL,
  geneSymbol = NULL,
  main = NULL,
  probe2gene = TRUE,
  ...
)
```

## Arguments

- |            |  |
|------------|--|
| object     | ExpressionSet object for the experiment  |
| groups     | String containing the name of the grouping variable. This should be a the name of a column in the pData of the expressionSet object. |
| probesetId | The probeset ID. These should be stored in the featureNames of the expressionSet object.   |

geneSymbol	The gene symbol. These should be stored in the column `Gene Symbol` in the featureData of the expressionSet object.
main	Main title on top of the graph
probe2gene	Boolean indicating whether the probeset should be translated to a gene symbol (used for the default title of the plot)
...	Possibility to add extra plot options. See <a href="#">par</a>

**Value**

a plot is drawn in the current device. prediction object is returned invisibly.

**Author(s)**

Willem Talloen

**References**

Some explanation about ROC can be found on [http://en.wikipedia.org/wiki/ROC\\_curve](http://en.wikipedia.org/wiki/ROC_curve) and <http://www.anaesthetist.com/mnm/stats/roc/Findex.htm>. The latter has at the bottom a nice interactive tool to scroll the cut-off and to see how it affects the FP/TP table and the ROC curve.

**Examples**

```
# simulated data set
esSim <- simulateData()
ROCCurve(probesetId = 'Gene.1', object = esSim, groups = 'type', addLegend = FALSE)
```

**topTable,pamClass-method**

*Top table for pamClass object*

**Description**

Top table for pamClass object

**Usage**

```
## S4 method for signature 'pamClass'
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

**Value**

`topTablePam` object

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`topTable, rfClass-method`

*Top table for rfClass object*

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**Description**

Top table for `rfClass` object

**Usage**

```
## S4 method for signature 'rfClass'  
topTable(fit, n)
```

**Arguments**

<code>fit</code>	object for which to obtain a top table, generally a fit object for a given model class
<code>n</code>	number of features (variables) to list in the top table, ranked by importance

**Value**

`topTableRfClass` object

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