

# Package ‘ClassifyR’

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**Type** Package

**Title** A framework for cross-validated classification problems, with applications to differential variability and differential distribution testing

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**Suggests** limma, genefilter, edgeR, car, Rmixmod, ggplot2 (>= 3.0.0), gridExtra (>= 2.0.0), cowplot, BiocStyle, pamr, PoiClaClu, parathyroidSE, knitr, htmltools, gtable, scales, e1071, rmarkdown, IRanges, randomForest, robustbase, glmnet, class

**Description** The software formalises a framework for classification in R. There are four stages; Data transformation, feature selection, classifier training, and prediction. The requirements of variable types and names are fixed, but specialised variables for functions can also be provided. The classification framework is wrapped in a driver loop, that reproducibly carries out a number of cross-validation schemes. Functions for differential expression, differential variability, and differential distribution are included. Additional functions may be developed by the user, by creating an interface to the framework.

**Collate** classes.R utilities.R bartlettSelection.R calcPerformance.R classifyInterface.R DLDAinterface.R DMDselection.R differentMeansSelection.R distribution.R easyHardClassifier.R easyHardFeatures.R edgeRselection.R edgesToHubNetworks.R elasticNetGLMinterface.R elasticNetFeatures.R featureSetSummary.R fisherDiscriminant.R forestFeatures.R

getLocationAndScales.R interactorDifferences.R kNNinterface.R  
 KolmogorovSmirnovSelection.R KullbackLeiblerSelection.R  
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 limmaSelection.R mixmodels.R naiveBayesKernel.R  
 networkCorrelationsSelection.R NSCselectionInterface.R  
 NSCtrainInterface.R NSCpredictInterface.R  
 pairsDifferencesSelection.R performancePlot.R  
 plotFeatureClasses.R previousSelection.R previousTrained.R  
 randomForestInterface.R rankingPlot.R ROCplot.R runTest.R  
 runTests.R samplesMetricMap.R selectionPlot.R  
 subtractFromLocation.R SVMinterface.R

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asthma

*Asthma RNA Abundance and Patient Classes*

---

### Description

Data set consists of a matrix of abundances of 2000 most variable gene expression measurements for 190 samples and a factor vector of classes for those samples.

### Usage

```
data(asthma)
```

### Format

measurements has a row for each gene and a column for each sample. classes is a factor vector.

### Source

A Nasal Brush-based Classifier of Asthma Identified by Machine Learning Analysis of Nasal RNA Sequence Data, *Scientific Reports*, 2018. Webpage: <http://www.nature.com/articles/s41598-018-27189-4>

---

bartlettSelection

*Selection of Differential Variability with Bartlett Statistic*

---

### Description

Ranks features by largest Bartlett statistic and chooses the features which have best resubstitution performance.

### Usage

```
## S4 method for signature 'matrix'
bartlettSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
bartlettSelection(measurements, classes, datasetName,
                  trainParams, predictParams, resubstituteParams,
                  selectionName = "Bartlett Test", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
bartlettSelection(measurements, targets, ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the matrix nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

The calculation of the test statistic is performed by the [bartlett.test](#) function from the [stats](#) package.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class [MultiAssayExperiment](#), the factor of sample classes must be stored in the [DataFrame](#) accessible by the [colData](#) function with column name "class".

**Value**

An object of class [SelectResult](#) or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

**Author(s)**

Dario Strbenac

**Examples**

```
# Samples in one class with differential variability to other class.
# First 20 genes are DV.
genesRNAmatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 1)))
moreVariable <- sapply(1:25, function(sample) rnorm(20, 9, 5))
genesRNAmatrix <- cbind(genesRNAmatrix, rbind(moreVariable,
```

```

      sapply(1:25, function(sample) rnorm(80, 9, 1)))
colnames(genesRNAmatrix) <- paste("Sample", 1:50)
rownames(genesRNAmatrix) <- paste("Gene", 1:100)
genesSNPmatrix <- matrix(sample(c("None", "Missense"), 250, replace = TRUE),
      ncol = 50)
colnames(genesSNPmatrix) <- paste("Sample", 1:50)
rownames(genesSNPmatrix) <- paste("Gene", 1:5)
classes <- factor(rep(c("Poor", "Good"), each = 25))
names(classes) <- paste("Sample", 1:50)
genesDataset <- MultiAssayExperiment(list(RNA = genesRNAmatrix, SNP = genesSNPmatrix),
      colData = DataFrame(class = classes))
# Wait for update to MultiAssayExperiment wideFormat function.
trainIDs <- paste("Sample", c(1:20, 26:45))
genesDataset <- subtractFromLocation(genesDataset, training = trainIDs,
      targets = "RNA") # Exclude SNP data.

resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
      performanceType = "balanced error",
      better = "lower")
bartlettSelection(genesDataset, datasetName = "Example", targets = "RNA",
      trainParams = TrainParams(fisherDiscriminant),
      predictParams = PredictParams(NULL),
      resubstituteParams = resubstituteParams)

```

---

calcPerformance	<i>Add Performance Calculations to a ClassifyResult Object or Calculate for a Pair of Factor Vectors</i>
-----------------	--

---

## Description

If `calcExternalPerformance` is used, such as when having a vector of known classes and a vector of predicted classes determined outside of the `ClassifyR` package, a single metric value is calculated. If `calcCVperformance` is used, annotates the results of calling `runTests` with one of the user-specified performance measures.

## Usage

```

## S4 method for signature 'factor,factor'
calcExternalPerformance(actualClasses, predictedClasses,
      performanceType = c("error", "accuracy", "balanced error", "balanced accuracy",
        "micro precision", "micro recall",
        "micro F1", "macro precision",
        "macro recall", "macro F1", "matthews"))

## S4 method for signature 'ClassifyResult'
calcCVperformance(result, performanceType = c("error", "accuracy", "balanced error", "balanced accuracy",
      "sample error", "sample accuracy",
      "micro precision", "micro recall",
      "micro F1", "macro precision",
      "macro recall", "macro F1", "matthews"))

```

## Arguments

result	An object of class <code>ClassifyResult</code> .
performanceType	<p>A character vector of length 1. Default: "balanced error". Must be one of the following options:</p> <ul style="list-style-type: none"> <li>• "error": Ordinary error rate.</li> <li>• "accuracy": Ordinary accuracy.</li> <li>• "balanced error": Balanced error rate.</li> <li>• "balanced accuracy": Balanced accuracy.</li> <li>• "sample error": Error rate for each sample in the data set.</li> <li>• "sample accuracy": Accuracy for each sample in the data set.</li> <li>• "micro precision": Sum of the number of correct predictions in each class, divided by the sum of number of samples in each class.</li> <li>• "micro recall": Sum of the number of correct predictions in each class, divided by the sum of number of samples predicted as belonging to each class.</li> <li>• "micro F1": F1 score obtained by calculating the harmonic mean of micro precision and micro recall.</li> <li>• "macro precision": Sum of the ratios of the number of correct predictions in each class to the number of samples in each class, divided by the number of classes.</li> <li>• "macro recall": Sum of the ratios of the number of correct predictions in each class to the number of samples predicted to be in each class, divided by the number of classes.</li> <li>• "macro F1": F1 score obtained by calculating the harmonic mean of macro precision and macro recall.</li> <li>• "matthews": Matthews Correlation Coefficient (MCC). A score between -1 and 1 indicating how concordant the predicted classes are to the actual classes. Only defined if there are two classes.</li> </ul>
actualClasses	A factor vector specifying each sample's correct class.
predictedClasses	A factor vector of the same length as actualClasses specifying each sample's predicted class.

## Details

All metrics except Matthews Correlation Coefficient are suitable for evaluating classification scenarios with more than two classes and are reimplementations of those available from [Intel DAAL](#).

If `runTests` was run in resampling mode, one performance measure is produced for every resampling. If the leave-k-out mode was used, then the predictions are concatenated, and one performance measure is calculated for all classifications.

"balanced error" calculates the balanced error rate and is better suited to class-imbalanced data sets than the ordinary error rate specified by "error". "sample error" calculates the error rate of each sample individually. This may help to identify which samples are contributing the most to the

overall error rate and check them for confounding factors. Precision, recall and F1 score have micro and macro summary versions. The macro versions are preferable because the metric will not have a good score if there is substantial class imbalance and the classifier predicts all samples as belonging to the majority class.

### Value

If `calcCVperformance` was run, an updated `ClassifyResult` object, with new metric values in the performance slot. If `calcExternalPerformance` was run, the performance metric value itself.

### Author(s)

Dario Strbenac

### Examples

```
predictTable <- data.frame(sample = paste("A", 1:10, sep = ''),
                           class = factor(sample(LETTERS[1:2], 50, replace = TRUE)))
actual <- factor(sample(LETTERS[1:2], 10, replace = TRUE))
result <- ClassifyResult("Example", "Differential Expression", "A Selection",
                        paste("A", 1:10, sep = ''), paste("Gene", 1:50, sep = ''),
                        50, list(1:50, 1:50), list(1:5, 6:15), list(function(oracle){}),
                        list(predictTable), actual, list("leave", 2))
result <- calcCVperformance(result, "balanced error")
performance(result)
```

---

characterOrDataFrame *Union of a Character and a DataFrame*

---

### Description

Allows a slot to be either a character or a DataFrame.

### Author(s)

Dario Strbenac

### Examples

```
setClass("Selections", representation(features = "characterOrDataFrame"))
selections <- new("Selections", features = c("BRAF", "NRAS"))
featuresTable <- DataFrame(assay = c("RNA-seq", "Mass spectrometry"),
                          feature = c("PD-1", "MITF"))
omicsSelections <- new("Selections", features = featuresTable)
```



---

classifyInterface      *An Interface for PoiClaClu Package's Classify Function*

---

### Description

More details of Poisson LDA are available in the documentation of [Classify](#).

### Usage

```
## S4 method for signature 'matrix'
classifyInterface(measurements, classes, test, ...)
## S4 method for signature 'DataFrame'
classifyInterface(measurements, classes, test, ..., returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
classifyInterface(measurements, test, targets = names(measurements), ...)
```

### Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples. If of type <a href="#">DataFrame</a> , the data set is subset to only those features of type integer.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or parameters that <a href="#">Classify</a> can accept.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, matrix of scores for each class, or both labels and scores in a data.frame.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

Data tables which consist entirely of non-integer data cannot be analysed. If measurements is an object of class [MultiAssayExperiment](#), the factor of sample classes must be stored in the [DataFrame](#) accessible by the `colData` function with column name "class".

**Value**

Either a factor vector of predicted classes, a matrix of scores for each class, or a table of both the class labels and class scores, depending on the setting of `returnType`.

**Author(s)**

Dario Strbenac

**Examples**

```
if(require(PoiClaClu))
{
  readCounts <- CountDataSet(n = 100, p = 1000, 2, 5, 0.1)
  # Rows are for features, columns are for samples.
  trainData <- t(readCounts[['x']])
  classes <- factor(paste("Class", readCounts[['y']]))
  testData <- t(readCounts[['xte']])
  storage.mode(trainData) <- storage.mode(testData) <- "integer"
  classified <- classifyInterface(trainData, classes, testData)

  setNames(table(paste("Class", readCounts[['yte']]) == classified), c("Incorrect", "Correct"))
}
```

---

ClassifyResult

*Container for Storing Classification Results*

---

**Description**

Contains a table of actual sample classes and predicted classes, the identifiers of features selected for each fold of each permutation or each hold-out classification, and error rates. This class is not intended to be created by the user, but could be used in another package. It is created by [runTests](#).

**Constructor**

```
ClassifyResult(datasetName, classificationName, selectionName, originalNames, originalFeatures,
               rankedFeatures, chosenFeatures, predictions, actualClasses, models,
               validation, tune = list(NULL))
```

`datasetName` A name associated with the dataset used.

`classificationName` A name associated with the classification.

`selectionName` A name associated with the feature selection.

`originalNames` All sample names.

`originalFeatures` All feature names. Character vector or `DataFrame` with one row for each feature if the data set is a [MultiAssayExperiment](#).

`rankedFeatures` All features, from most to least important. Character vector or `DataFrame` if data set is a `MultiAssayExperiment`.

`chosenFeatures` Features selected at each fold. Character vector or `DataFrame` if data set is a `MultiAssayExperiment`.

`predictions` A `list` of `data.frame` containing information about samples, their actual class and predicted class.

`actualClasses` Factor of class of each sample.

`models` All of the models fitted to the training data.

`validation` List with first element being the name of the validation scheme, and other elements providing details about the scheme.

`tune` A description of the tuning parameters, and the value chosen of each parameter.

### Summary

A method which summarises the results is available. `result` is a `ClassifyResult` object.

`show(result)` Prints a short summary of what `result` contains.

`totalPredictions(ClassifyResult)` Calculates the sum of the number of predictions.

### Accessors

`result` is a `ClassifyResult` object.

`sampleNames(result)` Returns a vector of sample names present in the data set.

`featureNames(result)` Returns a vector of features present in the data set.

`predictions(result)` Returns a `list` of `data.frame`. Each `data.frame` contains columns `sample`, `predicted`, and `actual`. For hold-out validation, only one `data.frame` is returned of all of the concatenated predictions.

`actualClasses(result)` Returns a factor class labels, one for each sample.

`features(result)` A `list` of the features selected for each training.

`models(result)` A `list` of the models fitted for each training.

`performance(result)` Returns a `list` of performance measures. This is empty until `calcCVperformance` has been used.

`tunedParameters(result)` Returns a `list` of tuned parameter values. If cross-validation is used, this list will be large, as it stores chosen values for every iteration.

`sampleNames(result)` Returns a `character` vector of sample names.

### Author(s)

Dario Strbenac

**Examples**

```

#if(require(sparsediscrim))
#{
  data(asthma)

  resubstituteParams <- ResubstituteParams(nFeatures = seq(5, 25, 5),
                                           performanceType = "balanced error",
                                           better = "lower")

  classified <-
  runTests(measurements, classes, datasetName = "Asthma",
           classificationName = "Different Means", validation = "leaveOut", leave = 1,
           params = list(SelectParams(limmaSelection, "Moderated t Statistic",
                                     resubstituteParams = resubstituteParams),
                        TrainParams(DLDAtrainInterface),
                        PredictParams(DLDApredictInterface)))

  class(classified)
#}

```

---

differentMeansSelection

*Selection of Differentially Abundant Features*


---

**Description**

Uses an ordinary t-test if the data set has two classes or one-way ANOVA if the data set has three or more classes to select differentially expressed features.

**Usage**

```

## S4 method for signature 'matrix'
differentMeansSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
differentMeansSelection(measurements, classes, datasetName,
                       trainParams, predictParams, resubstituteParams,
                       selectionName = "Difference in Means", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
differentMeansSelection(measurements, targets = NULL, ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	A vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	Names of data tables to be combined into a single table and used in the analysis.

...	Variables not used by the <code>matrix</code> nor the <code>MultiAssayExperiment</code> method which are passed into and used by the <code>DataFrame</code> method.
<code>datasetName</code>	A name for the data set used. Stored in the result.
<code>trainParams</code>	A container of class <code>TrainParams</code> describing the classifier to use for training.
<code>predictParams</code>	A container of class <code>PredictParams</code> describing how prediction is to be done.
<code>resubstituteParams</code>	An object of class <code>ResubstituteParams</code> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
<code>selectionName</code>	A name to identify this selection method by. Stored in the result.
<code>verbose</code>	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

This selection method looks for changes in means and uses `rowttests` to rank the features if there are two classes or `rowFtests` if there are three or more classes. The choice of features is based on the best resubstitution performance.

### Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac

### Examples

```
#if(require(sparsediscrim))
#{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))

  resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "balanced error", better = "lower")
  selected <- differentMeansSelection(genesMatrix, classes, "Example",
    trainParams = TrainParams(), predictParams = PredictParams(),
    resubstituteParams = resubstituteParams)

  selected@chosenFeatures
#}
```

---

distribution

*Get Frequencies of Feature Selection and Sample Errors*


---

## Description

There are two modes. For aggregating feature selection results, the function counts the number of times each feature was selected in all cross-validations. For aggregating classification results, the error rate for each sample is calculated. This is useful in identifying outlier samples that are difficult to classify.

## Usage

```
## S4 method for signature 'ClassifyResult'
distribution(result, dataType = c("features", "samples"),
            plotType = c("density", "histogram"), summaryType = c("percentage", "count"),
            plot = TRUE, xMax = NULL, xLabel = "Percentage of Cross-validations",
            yLabel = "Density", title = "Distribution of Feature Selections",
            fontSizes = c(24, 16, 12), ...)
```

## Arguments

result	An object of class <a href="#">ClassifyResult</a> .
dataType	Whether to calculate sample-wise error rate or the number of times a feature was selected.
plotType	Whether to draw a probability density curve or a histogram.
summaryType	Whether to summarise the feature selections as a percentage or count.
plot	Whether to draw a plot of the frequency of selection or error rate.
xMax	Maximum data value to show in plot.
xLabel	The label for the x-axis of the plot.
yLabel	The label for the y-axis of the plot.
title	An overall title for the plot.
fontSizes	A vector of length 3. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values.
...	Further parameters, such as colour and fill, passed to <a href="#">geom_histogram</a> or <a href="#">stat_density</a> , depending on the value of plotType.

## Value

If type is "features", a vector as long as the number of features that were chosen at least once containing the number of times the feature was chosen in cross validations or the percentage of times chosen. If type is "samples", a vector as long as the number of samples, containing the cross-validation error rate of the sample. If plot is TRUE, then a plot is also made on the current graphics device.

**Author(s)**

Dario Strbenac

**Examples**

```

#if(require(sparsediscrim))
#{
  data(asthma)
  resubstituteParams <- ResubstituteParams(nFeatures = seq(5, 25, 5),
                                          performanceType = "balanced error",
                                          better = "lower")

  result <- runTests(measurements, classes, datasetName = "Asthma",
                    classificationName = "Different Means",
                    permutations = 5,
                    params = list(SelectParams(limmaSelection, "Moderated t Statistic",
                                              resubstituteParams = resubstituteParams),
                                  TrainParams(DLDAtrainInterface),
                                  PredictParams(DLDApredictInterface)
                    )
  )
  featureDistribution <- distribution(result, "features", summaryType = "count",
                                    plotType = "histogram",
                                    xLabel = "Number of Cross-validations", yLabel = "Count",
                                    binwidth = 1)
  print(head(featureDistribution))
#}

```

---

*dlda**Trained dlda Object*

---

**Description**

Enables S4 method dispatching on it.

**Author(s)**

Dario Strbenac

---

*DLDAinterface**An Interface for sparsediscrim Package's dlda Function*

---

**Description**

DLDAtrainInterface generates a trained diagonal LDA classifier and DLDApredictInterface uses it to make predictions on a test data set.

**Usage**

```

## S4 method for signature 'matrix'
DLDAtrainInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
DLDAtrainInterface(measurements, classes, verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
DLDAtrainInterface(measurements, targets = names(measurements), ...)
## S4 method for signature 'dllda,matrix'
DLDApredictInterface(model, test, ...)
## S4 method for signature 'dllda,DataFrame'
DLDApredictInterface(model, test, returnType = c("both", "class", "score"),
                      verbose = 3)
## S4 method for signature 'dllda,MultiAssayExperiment'
DLDApredictInterface(model, test, targets = names(test), ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples. If of type <a href="#">DataFrame</a> , the data set is subset to only those features of type integer.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
model	A fitted model as returned by <a href="#">DLDAtrainInterface</a> .
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it. Also, if a <a href="#">DataFrame</a> , the class column must be absent.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method (e.g. <code>verbose</code> ).
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, matrix of scores for each class, or both labels and scores in a data.frame.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

If `measurements` is an object of class [MultiAssayExperiment](#), the factor of sample classes must be stored in the [DataFrame](#) accessible by the `colData` function with column name "class".



**Value**

For `DLDAtrainInterface`, a trained DLDA classifier. For `DLDApredictInterface`, either a factor vector of predicted classes, a matrix of scores for each class, or a table of both the class labels and class scores, depending on the setting of `returnType`.

**Author(s)**

Dario Strbenac

**Examples**

```
# if(require(sparsediscrim)) Package currently removed from CRAN.
#{
# Genes 76 to 100 have differential expression.
genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
  c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
classes <- factor(rep(c("Poor", "Good"), each = 25))
colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
selected <- rownames(genesMatrix)[91:100]
trainingSamples <- c(1:20, 26:45)
testingSamples <- c(21:25, 46:50)

classifier <- DLDAtrainInterface(genesMatrix[selected, trainingSamples],
  classes[trainingSamples])
DLDApredictInterface(classifier, genesMatrix[selected, testingSamples])
#}
```

---

DMDselection

*Selection of Differential Distributions with Differences in Means or Medians and a Deviation Measure*

---

**Description**

Ranks features by largest Differences in Means/Medians and Deviations and chooses the features which have best resubstitution performance.

**Usage**

```
## S4 method for signature 'matrix'
DMDselection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
DMDselection(measurements, classes, datasetName, differences = c("both", "location", "scale"),
  trainParams, predictParams, resubstituteParams, ...,
  selectionName = "Differences of Medians and Deviations",
  verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
DMDselection(measurements, targets = names(measurements), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	A vector of class labels.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or parameters for <a href="#">getLocationsAndScales</a> , such as <a href="#">location</a> , <a href="#">scale</a> .
datasetName	Default: "Differences of Medians and Deviations". A name for the data set used. Stored in the result.
differences	Default: "both". Either "both", "location", or "scale". The type of differences to consider. If both are considered then the absolute difference in location and the absolute difference in scale are summed.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

DMD is defined as  $\sum_{i=1} \sum_{j=i+1} |location_i - location_j| + |scale_i - scale_j|$ . The subscripts denote the class for which the parameter is calculated for.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class [MultiAssayExperiment](#), the factor of sample classes must be stored in the [DataFrame](#) accessible by the [colData](#) function with column name "class".

**Value**

An object of class [SelectResult](#) or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

**Author(s)**

Dario Strbenac

**Examples**

```
# First 20 features have bimodal distribution for Poor class.
# Other 80 features have normal distribution for both classes.
genesMatrix <- sapply(1:25, function(sample)
  {
    randomMeans <- sample(c(8, 12), 20, replace = TRUE)
    c(rnorm(20, randomMeans, 1), rnorm(80, 10, 1))
  }
)
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample) rnorm(100, 10, 1)))
classes <- factor(rep(c("Poor", "Good"), each = 25))

resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
  performanceType = "balanced error",
  better = "lower")
DMDselection(genesMatrix, classes, datasetName = "Example",
  trainParams = TrainParams(naiveBayesKernel),
  predictParams = PredictParams(NULL),
  resubstituteParams = resubstituteParams)
```

---

EasyHardClassifier      *Container for a Pair of Trained Classifiers*

---

**Description**

Stores two classifiers and the ID of the data set each one of them was trained on. Not intended for end-user.

**Constructor**

`EasyHardClassifier(easyClassifier, hardClassifier, datasetIDs)`

Creates a `EasyHardClassifier` object which stores the details of the two underlying classification models.

`easyClassifier` A classifier trained on the easy-to-collect data set by [easyHardClassifier-Train](#). Represented as a list of rules, each of which is also a list.

`hardClassifier` A list of two with names "selected" and "model". The "selected" element should contain the chosen features and the "model" element is for the trained model on the hard-to-collect data set or simply a character vector of length 1 containing a class name, if the prediction of the samples left over from the easy classifier all or all except one belonged to a particular class.

`datasetIDs` A vector of length 2 with names "easy" and "hard" containing the data set IDs from the original [MultiAssayExperiment](#) input data object for the easy data set and the hard data set.

**Author(s)**

Dario Strbenac

## Examples

```
predictiveRules <- list(list(feature = "age", relation = "<", value = 18, predict = "Risk"))
hardClassifier <- DLDAtrainInterface(matrix(rnorm(400), ncol = 20),
  classes = factor(sample(c("Safe", "Risk"), 20, replace = TRUE)))
EasyHardClassifier(predictiveRules, list(selected = LETTERS[1:5], model = hardClassifier),
  setNames(c("clinical", "RNA-seq"), c("easy", "hard")))
```

---

easyHardClassifier	<i>Two-stage Classification Using Easy-to-collect Data Set and Hard-to-collect data set.</i>
--------------------	--

---

## Description

An alternative implementation to the previously published easy-hard classifier that doesn't do nested cross-validation for speed. In the first stage, each numeric variable is split on all possible midpoints between consecutive ordered values and the samples below the split and above the split are checked to see if they mostly belong to one class. Categorical variables are tabulated on factor levels and the count of samples in each class is determined. If any partitions of samples are pure for a class, based on a purity threshold, prediction rules are created. The samples not classified by any rule or classified to two or more classes the same number of times are left to be trained by the hard classifier.

## Usage

```
## S4 method for signature 'MultiAssayExperiment'
easyHardClassifierTrain(measurements, easyDatasetID = "clinical", hardDatasetID = names(measurements)
  featureSets = NULL, metaFeatures = NULL, minimumOverlapPercent = 80,
  datasetName = NULL, classificationName = "Easy-Hard Classifier",
  easyClassifierParams = list(minCardinality = 10, minPurity = 0.9),
  hardClassifierParams = list(SelectParams(), TrainParams(), PredictParams()),
  verbose = 3)
## S4 method for signature 'EasyHardClassifier,MultiAssayExperiment'
easyHardClassifierPredict(model, test, predictParams, verbose = 3)
```

## Arguments

measurements	A <a href="#">MultiAssayExperiment</a> object containing the data set The sample classes must be in a column of the DataFrame accessed by colData named "class".
easyDatasetID	The name of a data set in measurements or "clinical" to indicate the patient information in the column data be used.
hardDatasetID	The name of a data set in measurements different to the value of easyDatasetID to be used for classifying the samples not classified by the easy classifier.
featureSets	An object of type <a href="#">FeatureSetCollection</a> which defines sets of features or sets of edges.
metaFeatures	Either NULL or a DataFrame which has meta-features of the numeric data of interest.

minimumOverlapPercent	If featureSets stores sets of features, the minimum overlap of feature IDs with measurements for a feature set to be retained in the analysis. If featureSets stores sets of network edges, the minimum percentage of edges with both vertex IDs found in measurements that a set has to have to be retained in the analysis.
datasetName	A name associated with the data set used.
classificationName	A name associated with the classification.
easyClassifierParams	A list of length 2 with names "minCardinality" and "minPurity". The first parameter specifies what the minimum number of samples after a split has to be and the second specifies the minimum proportion of samples in a partition belonging to a particular class.
hardClassifierParams	A list of objects defining the classification to do on the samples which were not predicted by the easy classifier Objects must of of class <a href="#">TransformParams</a> , <a href="#">SelectParams</a> , <a href="#">TrainParams</a> or <a href="#">PredictParams</a> .
model	A trained <a href="#">EasyHardClassifier</a> object.
test	A <a href="#">MultiAssayExperiment</a> object containing the test data.
predictParams	An object of class <a href="#">PredictParams</a> . It specifies the classifier used to make the hard predictions.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

The easy classifier may be NULL if there are no rules that predicted the sample well using the easy data set. The hard classifier may be NULL if all of the samples could be predicted with rules generated using the easy data set or it will simply be a character if all or almost all of the remaining samples belong to one class.

### Value

For `EasyHardClassifierTrain`, the trained two-stage classifier. For `EasyHardClassifierPredict`, a factor vector of predicted classes.

### Author(s)

Dario Strbenac

### References

Inspired by: Stepwise Classification of Cancer Samples Using Clinical and Molecular Data, Askar Obulkasim, Gerrit Meijer and Mark van de Wiel 2011, *BMC Bioinformatics*, Volume 12 article 422, <https://bmcbioinformatics.biomedcentral.com/articles/10.1186/1471-2105-12-422>.

## Examples

```

genesMatrix <- matrix(c(rnorm(90, 9, 1),
                       9.5, 9.4, 5.2, 5.3, 5.4, 9.4, 9.6, 9.9, 9.1, 9.8),
                     ncol = 10, byrow = TRUE)
colnames(genesMatrix) <- paste("Sample", 1:10)
rownames(genesMatrix) <- paste("Gene", 1:10)
genders <- factor(c("Male", "Male", "Female", "Female", "Female",
                   "Female", "Female", "Female", "Female", "Female"))

# Scenario: Male gender can predict the hard-to-classify Sample 1 and Sample 2.
clinical <- DataFrame(age = c(31, 34, 32, 39, 33, 38, 34, 37, 35, 36),
                     gender = genders,
                     class = factor(rep(c("Poor", "Good"), each = 5)),
                     row.names = colnames(genesMatrix))
dataset <- MultiAssayExperiment(ExperimentList(RNA = genesMatrix), clinical)
selParams <- SelectParams(featureSelection = differentMeansSelection, selectionName = "Difference in Means",
                          resubstituteParams = ResubstituteParams(1:10, "balanced error", "lower"))
trained <- easyHardClassifierTrain(dataset, easyClassifierParams = list(minCardinality = 2, minPurity = 0.9),
                                  hardClassifierParams = list(selParams, TrainParams(), PredictParams()))

predictions <- easyHardClassifierPredict(trained, dataset, PredictParams())

```

---

easyHardFeatures

*Extract Chosen Features from an EasyHardClassifier Object*

---

## Description

The features are described by a data frame. One column is named "dataset" and the other is named "feature". This provides identifiability in case when multiple types of data have features with the same name.

## Usage

```

## S4 method for signature 'EasyHardClassifier'
easyHardFeatures(easyHardClassifier)

```

## Arguments

easyHardClassifier  
 An [EasyHardClassifier](#) object.

## Details

If any of the features are from the column data of the input [MultiAssayExperiment](#), the dataset value will be "clinical".

**Value**

To be consistent with other functions for extracting features from a trained model, a list of length two. The first element is for feature rankings, which is not meaningful for an easy-hard classifier, so it is NULL. The second element is the selected features.

**Author(s)**

Dario Strbenac

**Examples**

```
genesMatrix <- matrix(c(rnorm(90, 9, 1),
                        9.5, 9.4, 5.2, 5.3, 5.4, 9.4, 9.6, 9.9, 9.1, 9.8),
                      ncol = 10, byrow = TRUE)
colnames(genesMatrix) <- paste("Sample", 1:10)
rownames(genesMatrix) <- paste("Gene", 1:10)
genders <- factor(c("Male", "Male", "Female", "Female", "Female",
                   "Female", "Female", "Female", "Female", "Female"))

# Scenario: Male gender can predict the hard-to-classify Sample 1.
clinical <- DataFrame(age = c(31, 34, 32, 39, 33, 38, 34, 37, 35, 36),
                     gender = genders,
                     class = factor(rep(c("Poor", "Good"), each = 5)),
                     row.names = colnames(genesMatrix))
dataset <- MultiAssayExperiment(ExperimentList(RNA = genesMatrix), clinical)
trained <- easyHardClassifierTrain(dataset, easyClassifierParams = list(minCardinality = 2, minPurity = 0.9),
                                hardClassifierParams = list(SelectParams(featureSelection = differentMeansSelection,
                                selectionName = "Difference in Means",
                                resubstituteParams = ResubstituteParams(1:10, "balanced error", "low

easyHardFeatures(trained)
```

---

edgeRselection

*Feature Selection Based on Differential Expression for Count Data*

---

**Description**

Performs a differential expression analysis between classes and chooses the features which have best resubstitution performance. The data may have overdispersion and this is modelled.

**Usage**

```
## S4 method for signature 'matrix'
edgeRselection(counts, classes, ...)
## S4 method for signature 'DataFrame'
edgeRselection(counts, classes, datasetName,
               normFactorsOptions = NULL, dispOptions = NULL, fitOptions = NULL,
               trainParams, predictParams, resubstituteParams,
```

```

        selectionName = "edgeR LRT", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
edgeRselection(counts, targets = NULL, ...)

```

### Arguments

counts	Either a <a href="#">matrix</a> or <a href="#">MultiAssayExperiment</a> containing the unnormalised counts.
classes	A vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables of counts to be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
datasetName	A name for the data set used. Stored in the result.
normFactorsOptions	A named list of any options to be passed to <a href="#">calcNormFactors</a> .
dispOptions	A named list of any options to be passed to <a href="#">estimateDisp</a> .
fitOptions	A named list of any options to be passed to <a href="#">glmFit</a> .
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

The differential expression analysis follows the standard [edgeR](#) steps of estimating library size normalisation factors, calculating dispersion, in this case robustly, and then fitting a generalised linear model followed by a likelihood ratio test.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class [MultiAssayExperiment](#), the factor of sample classes must be stored in the [DataFrame](#) accessible by the [colData](#) function with column name "class".

### Value

An object of class [SelectResult](#) or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac



## References

edgeR: a Bioconductor package for differential expression analysis of digital gene expression data, Mark D. Robinson, Davis McCarthy, and Gordon Smyth, 2010, *Bioinformatics*, Volume 26 Issue 1, <https://academic.oup.com/bioinformatics/article/26/1/139/182458>.

## Examples

```
if(require(parathyroidSE) && require(PoiClaClu))
{
  data(parathyroidGenesSE)
  expression <- assays(parathyroidGenesSE)[[1]]
  sampleNames <- paste("Sample", 1:ncol(parathyroidGenesSE))
  colnames(expression) <- sampleNames
  DPN <- which(colData(parathyroidGenesSE)[, "treatment"] == "DPN")
  control <- which(colData(parathyroidGenesSE)[, "treatment"] == "Control")
  expression <- expression[, c(control, DPN)]
  classes <- factor(rep(c("Control", "DPN"), c(length(control), length(DPN))))
  expression <- expression[rowSums(expression > 1000) > 8, ] # Make small data set.

  selected <- edgeRselection(expression, classes, "DPN Treatment",
    trainParams = TrainParams(classifyInterface),
    predictParams = PredictParams(NULL),
    resubstituteParams = ResubstituteParams(nFeatures = seq(10, 100, 10),
      performanceType = "balanced error", better = "lower"))

  head(selected@rankedFeatures[[1]])
  plotFeatureClasses(expression, classes, "ENSG00000044574",
    dotBinWidth = 500, xAxisLabel = "Unnormalised Counts")
}
```

---

edgesToHubNetworks      *Convert a Two-column Matrix or Data Frame into a Hub Node List*

---

## Description

Interactions between pairs of features (typically a protein-protein interaction, commonly abbreviated as PPI, database) are restructured into a named list. The name of the each element of the list is a feature and the element contains all features which have an interaction with it.

## Usage

```
edgesToHubNetworks(edges, minCardinality = 5)
```

## Arguments

**edges**      A two-column matrix or data.frame for which each row specifies a known interaction between two interactors. If feature X appears in the first column and feature Y appears in the second, there is no need for feature Y to appear in the first column and feature X in the second.

`minCardinality` An integer specifying the minimum number of features to be associated with a hub feature for it to be present in the result.

### Value

An object of type `FeatureSetCollection`.

### Author(s)

Dario Strbenac

### References

VAN: an R package for identifying biologically perturbed networks via differential variability analysis, Vivek Jayaswal, Sarah-Jane Schramm, Graham J Mann, Marc R Wilkins and Yee Hwa Yang, 2010, *BMC Research Notes*, Volume 6 Article 430, <https://bmcrnotes.biomedcentral.com/articles/10.1186/1756-0500-6-430>.

### Examples

```
interactor <- c("MITF", "MITF", "MITF", "MITF", "MITF", "MITF",
               "KRAS", "KRAS", "KRAS", "KRAS", "KRAS", "KRAS",
               "PD-1")
otherInteractor <- c("HINT1", "LEF1", "PSMD14", "PIAS3", "UBE2I", "PATZ1",
                    "ARAF", "CALM1", "CALM2", "CALM3", "RAF1", "HNRNPC",
                    "PD-L1")
edges <- data.frame(interactor, otherInteractor, stringsAsFactors = FALSE)

edgesToHubNetworks(edges, minCardinality = 4)
```

---

elasticNetFeatures	<i>Extract Vectors of Ranked and Selected Features From an Elastic Net GLM Object</i>
--------------------	---

---

### Description

Provides a ranking of features based on the magnitude of fitted GLM coefficients. Also provides the selected features which are those with a non-zero coefficient.

### Usage

```
## S4 method for signature 'multnet'
elasticNetFeatures(model)
```

### Arguments

`model` A fitted multinomial GLM which was created by `glmnet`.

**Value**

An list object. The first element is a vector or data frame of ranked features, the second is a vector or data frame of selected features.

**Author(s)**

Dario Strbenac

**Examples**

```

if(require(glmnet))
{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))

  resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "balanced error",
    better = "lower")

  # alpha is a user-specified tuning parameter.
  # lambda is automatically tuned, based on glmnet defaults, if not user-specified.
  trainParams <- TrainParams(elasticNetGLMtrainInterface, nlambda = 500)
  predictParams <- PredictParams(elasticNetGLMpredictInterface)
  classified <- runTests(genesMatrix, classes, datasetName = "Example",
    classificationName = "Differential Expression",
    validation = "fold",
    params = list(trainParams, predictParams))

  elasticNetFeatures(models(classified)[[1]])
}

```

---

elasticNetGLMinterface

*An Interface for glmnet Package's glmnet Function*

---

**Description**

An elastic net GLM classifier uses a penalty which is a combination of a lasso penalty and a ridge penalty, scaled by a lambda value, to fit a sparse linear model to the data.

**Usage**

```

## S4 method for signature 'matrix'
elasticNetGLMtrainInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
elasticNetGLMtrainInterface(measurements, classes, lambda = NULL,
                             ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
elasticNetGLMtrainInterface(measurements, targets = names(measurements), ...)
## S4 method for signature 'multnet,matrix'
elasticNetGLMpredictInterface(model, test, ...)
## S4 method for signature 'multnet,DataFrame'
elasticNetGLMpredictInterface(model, test, classes = NULL, lambda, ..., returnType = c("both", "class"))
## S4 method for signature 'multnet,MultiAssayExperiment'
elasticNetGLMpredictInterface(model, test, targets = names(test), ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples. If of type <a href="#">DataFrame</a> , the data set is subset to only those features of type integer.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
lambda	The lambda value passed directly to <a href="#">glmnet</a> if the training function is used or passed as s to <a href="#">predict.glmnet</a> if the prediction function is used.
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method (e.g. <code>verbose</code> ) or, for the training function, options that are used by the <a href="#">glmnet</a> function. For the testing function, this variable simply contains any parameters passed from the classification framework to it which aren't used by <a href="#">glmnet</a> 's <code>predict</code> function.
model	A trained elastic net GLM, as created by the <a href="#">glmnet</a> function.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, matrix of scores for each class, or both labels and scores in a <code>data.frame</code> .
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

If `measurements` is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name "class".

The value of the family parameter is fixed to "multinomial" so that classification with more than 2 classes is possible and type.multinomial is fixed to "grouped" so that a grouped lasso penalty is used. During classifier training, if more than one lambda value is considered by specifying a vector of them as input or leaving the default value of NULL, then the chosen value is determined based on classifier resubstitution error rate.

### Value

For elasticNetGLMtrainInterface, an object of type glmnet. For elasticNetGLMpredictInterface, either a factor vector of predicted classes, a matrix of scores for each class, or a table of both the class labels and class scores, depending on the setting of returnType.

### Author(s)

Dario Strbenac

### Examples

```
if(require(glmnet))
{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))

  resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "balanced error",
    better = "lower")

  # lambda is automatically tuned, based on glmnet defaults, if not user-specified.
  trainParams <- TrainParams(elasticNetGLMtrainInterface, nlambda = 500,
    getFeatures = elasticNetFeatures)
  predictParams <- PredictParams(elasticNetGLMpredictInterface)
  classified <- runTests(genesMatrix, classes, datasetName = "Example",
    classificationName = "Differential Expression",
    validation = "fold",
    params = list(trainParams, predictParams))

  classified <- calcCVperformance(classified, "balanced error")
  head(tunedParameters(classified))
  performance(classified)
}
```

---

FeatureSetCollection *Container for Storing A Collection of Sets*


---

**Description**

This container is the required storage format for a collection of sets. Typically, the elements of a set will either be a set of proteins (i.e. character vector) which perform a particular biological process or a set of binary interactions (i.e. Two-column matrix of feature identifiers).

**Constructor**

```
FeatureSetCollection(sets)
```

`sets` A named list. The names of the list describe the sets and the elements of the list specify the features which comprise the sets.

**Summary**

A method which summarises the results is available. `featureSets` is a `FeatureSetCollection` object.

```
show(featureSets)Prints a short summary of what featureSets contains.
```

**Subsetting**

The `FeatureSetCollection` may be subsetted or a single set may be extracted as a vector. `featureSets` is a `FeatureSetCollection` object.

```
featureSets[i:j]: Reduces the object to a subset of the feature sets between elements i and j of the collection.
```

```
featureSets[[i]]: Extract the feature set identified by i. i may be a numeric index or the character name of a feature set.
```

**Author(s)**

Dario Strbenac

**Examples**

```
ontology <- list(c("SES1", "PRDX1", "PRDX2", "PRDX3", "PRDX4", "PRDX5", "PRDX6",
  "LRRK2", "PARK7"),
  c("ATP7A", "CCS", "NQ01", "PARK7", "SOD1", "SOD2", "SOD3",
  "SZT2", "TNF"),
  c("AARS", "AIMP2", "CARS", "GARS", "KARS", "NARS", "NARS2",
  "LARS2", "NARS", "NARS2", "RGN", "UBA7"),
  c("CRY1", "CRY2", "ONP1SW", "OPN4", "RGR"),
  c("ESRRG", "RARA", "RARB", "RARG", "RXRA", "RXRB", "RXRG"),
  c("CD36", "CD47", "F2", "SDC4"),
  c("BUD31", "PARK7", "RWDD1", "TAF1")
```

```

)
names(ontology) <- c("Peroxioredoxin Activity", "Superoxide Dismutase Activity",
  "Ligase Activity", "Photoreceptor Activity",
  "Retinoic Acid Receptor Activity",
  "Thrombospondin Receptor Activity",
  "Regulation of Androgen Receptor Activity")

featureSets <- FeatureSetCollection(ontology)
featureSets
featureSets[3:5]
featureSets[["Photoreceptor Activity"]]

subNetworks <- list(MAPK = matrix(c("NRAS", "NRAS", "NRAS", "BRAF", "MEK",
  "ARAF", "BRAF", "CRAF", "MEK", "ERK"), ncol = 2),
  P53 = matrix(c("ATM", "ATR", "ATR", "P53",
  "CHK2", "CHK1", "P53", "MDM2"), ncol = 2)
)
networkSets <- FeatureSetCollection(subNetworks)
networkSets

```

---

FeatureSetCollectionOrNULL

*Union of a FeatureSetCollection and NULL*


---

### Description

Allows a slot to be either a FeatureSetCollectionOrNULL object or empty.

### Author(s)

Dario Strbenac

### Examples

```
TrainParams(DLDAtrainInterface, transform = NULL) # Use the input data as-is.
```

---

featureSetSummary

*Transform a Table of Feature Abundances into a Table of Feature Set Abundances.*


---

### Description

Represents a feature set by the mean or median feature measurement of a feature set for all features belonging to a feature set.

**Usage**

```
## S4 method for signature 'matrix'
featureSetSummary(measurements, location = c("median", "mean"),
                  featureSets, minimumOverlapPercent = 80, verbose = 3)
## S4 method for signature 'DataFrame'
featureSetSummary(measurements, location = c("median", "mean"),
                  featureSets, minimumOverlapPercent = 80, verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
featureSetSummary(measurements, target = NULL, location = c("median", "mean"),
                  featureSets, minimumOverlapPercent = 80, verbose = 3)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> or <a href="#">DataFrame</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
target	If the input is a <a href="#">MultiAssayExperiment</a> , this specifies which data set will be transformed. Can either be an integer or a character string specifying the name of the table. Must have length 1.
location	Default: The median. The type of location to summarise a set of features belonging to a feature set by.
featureSets	An object of type <a href="#">FeatureSetCollection</a> which defines the feature sets.
minimumOverlapPercent	The minimum percentage of overlapping features between the data set and a feature set defined in featureSets for that feature set to not be discarded from the analysis.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

This feature transformation method is unusual because the mean or median feature of a feature set for one sample may be different to another sample, whereas most other feature transformation methods do not result in different features being compared between samples during classification.

**Value**

The same class of variable as the input variable measurements is, with the individual features summarised to feature sets. The number of samples remains unchanged, so only one dimension of measurements is altered.

**Author(s)**

Dario Strbenac



## References

Network-based biomarkers enhance classical approaches to prognostic gene expression signatures, Rebecca L Barter, Sarah-Jane Schramm, Graham J Mann and Yee Hwa Yang, 2014, *BMC Systems Biology*, Volume 8 Supplement 4 Article S5, <https://bmcsystbiol.biomedcentral.com/articles/10.1186/1752-0509-8-S4-S5>.

## Examples

```
sets <- list(Adhesion = c("Gene 1", "Gene 2", "Gene 3"),
            `Cell Cycle` = c("Gene 8", "Gene 9", "Gene 10"))
featureSets <- FeatureSetCollection(sets)

# Adhesion genes have a median gene difference between classes.
genesMatrix <- matrix(c(rnorm(5, 9, 0.3), rnorm(5, 7, 0.3), rnorm(5, 8, 0.3),
                       rnorm(5, 6, 0.3), rnorm(10, 7, 0.3), rnorm(70, 5, 0.1)),
                     ncol = 10, byrow = TRUE)
rownames(genesMatrix) <- paste("Gene", 1:10)
colnames(genesMatrix) <- paste("Patient", 1:10)
classes <- factor(rep(c("Poor", "Good"), each = 5)) # But not used for transformation.

featureSetSummary(genesMatrix, featureSets = featureSets)
```

---

fisherDiscriminant      *Classification Using Fisher's LDA*

---

## Description

Finds the decision boundary using the training set, and gives predictions for the test set.

## Usage

```
## S4 method for signature 'matrix'
fisherDiscriminant(measurements, classes, test, ...)
## S4 method for signature 'DataFrame'
fisherDiscriminant(measurements, classes, test, returnType = c("both", "class", "score"),
                  verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
fisherDiscriminant(measurements, test, targets = names(measurements), ...)
```

## Arguments

**measurements**      Either a [matrix](#), [DataFrame](#) or [MultiAssayExperiment](#) containing the training data. For a matrix, the rows are features, and the columns are samples.

**classes**            Either a vector of class labels of class [factor](#) of the same length as the number of samples in **measurements** or if the measurements are of class [DataFrame](#) a character vector of length 1 containing the column name in measurement is also permitted. Not used if **measurements** is a [MultiAssayExperiment](#) object.

test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it.
targets	If measurements is a MultiAssayExperiment, the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the matrix nor the MultiAssayExperiment method which are passed into and used by the DataFrame method.
returnType	Default: "both". Either "class", "score", or "both". Sets the return value from the prediction to either a vector of class labels, score for a sample belonging to the second class, as determined by the factor levels, or both labels and scores in a data.frame.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

Unlike ordinary LDA, Fisher's version does not have assumptions about the normality of the features.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class MultiAssayExperiment, the factor of sample classes must be stored in the DataFrame accessible by the colData function with column name "class".

### Value

A vector or data.frame of class prediction information, as long as the number of samples in the test data.

### Author(s)

Dario Strbenac

### Examples

```
trainMatrix <- matrix(rnorm(1000, 8, 2), ncol = 10)
classes <- factor(rep(c("Poor", "Good"), each = 5))

# Make first 30 genes increased in value for poor samples.
trainMatrix[1:30, 1:5] <- trainMatrix[1:30, 1:5] + 5

testMatrix <- matrix(rnorm(1000, 8, 2), ncol = 10)

# Make first 30 genes increased in value for sixth to tenth samples.
testMatrix[1:30, 6:10] <- testMatrix[1:30, 6:10] + 5

fisherDiscriminant(trainMatrix, classes, testMatrix)
```

---

forestFeatures	<i>Extract Vectors of Ranked and Selected Features From a Random Forest Object</i>
----------------	--

---

### Description

Provides a ranking of features based on the total decrease in node impurities from splitting on the variable, averaged over all trees. Also provides the selected features which are those that were used in at least one tree of the forest.

### Usage

```
## S4 method for signature 'randomForest'  
forestFeatures(forest)
```

### Arguments

forest            A trained random forest which was created by [randomForest](#).

### Value

An list object. The first element is a vector or data frame of features, ranked from best to worst using the Gini index. The second element is a vector or data frame of features used in at least one tree.

### Author(s)

Dario Strbenac

### Examples

```
if(require(randomForest))  
{  
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))  
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)  
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))  
  classes <- factor(rep(c("Poor", "Good"), each = 25))  
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))  
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))  
  trainingSamples <- c(1:20, 26:45)  
  testingSamples <- c(21:25, 46:50)  
  
  trained <- randomForestTrainInterface(genesMatrix[, trainingSamples],  
    classes[trainingSamples], ntree = 10)  
  
  forestFeatures(trained)  
}
```

---

functionOrList      *Union of Functions and List of Functions*

---

**Description**

Allows a slot to be either a function or a list of functions.

**Author(s)**

Dario Strbenac

**Examples**

```
SelectParams(limmaSelection)
SelectParams(list(limmaSelection, leveneSelection), "Ensemble Selection")
```

---

functionOrNULL      *Union of A Function and NULL*

---

**Description**

Allows a slot to be either a function or empty.

**Author(s)**

Dario Strbenac

**Examples**

```
PredictParams(NULL)
PredictParams(predict)
```

---

getLocationsAndScales      *Calculate Location and Scale*

---

**Description**

Calculates the location and scale for each feature.

**Usage**

```
## S4 method for signature 'matrix'
getLocationsAndScales(measurements, ...)
## S4 method for signature 'DataFrame'
getLocationsAndScales(measurements, location = c("mean", "median"),
                      scale = c("SD", "MAD", "Qn"))
## S4 method for signature 'MultiAssayExperiment'
getLocationsAndScales(measurements, targets = names(measurements), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the data. For a matrix, the rows are features, and the columns are samples.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the matrix nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
location	The type of location to be calculated.
scale	The type of scale to be calculated.

**Details**

"SD" is used to represent standard deviation and "MAD" is used to represent median absolute deviation.

**Value**

A [list](#) of length 2. The first element contains the location for every feature. The second element contains the scale for every feature.

**Author(s)**

Dario Strbenac

**References**

Qn: <http://www.tandfonline.com/doi/pdf/10.1080/01621459.1993.10476408>

**Examples**

```
genesMatrix <- matrix(rnorm(1000, 8, 4), ncol = 10)
distributionInfo <- getLocationsAndScales(genesMatrix, "median", "MAD")

mean(distributionInfo[["median"]]) # Typical median.
mean(distributionInfo[["MAD"]]) # Typical MAD.
```

---

integerOrNumeric      *Union of a Integer and a Numeric*

---

### Description

Allows the same S4 subsetting function to be specified for object[i] and object[i:j], where i and j are integers.

### Author(s)

Dario Strbenac

### Examples

```
setClass("Container", representation(scores = "numeric"))
setMethod("[", c("Container", "integerOrNumeric", "missing", "ANY"),
  function(x, i, j, ..., drop = TRUE)
  {
    new("Container", scores = x@scores[i])
  })

dataset <- new("Container", scores = 1:10)
dataset[1] # 1 is numeric.
dataset[4:6] # 4:6 is a sequence of integers.
```

---

interactorDifferences      *Convert Individual Features into Differences Between Binary Interactors Based on Known Sub-networks*

---

### Description

This conversion is useful for creating a meta-feature table for classifier training and prediction based on sub-networks that were selected based on their differential correlation between classes.

### Usage

```
## S4 method for signature 'matrix'
interactorDifferences(measurements, ...)
## S4 method for signature 'DataFrame'
interactorDifferences(measurements, networkSets = NULL, absolute = FALSE, verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
interactorDifferences(measurements, target = NULL, ...)
```

**Arguments**

measurements	Either a <code>matrix</code> , <code>DataFrame</code> or <code>MultiAssayExperiment</code> containing the training data. For a <code>matrix</code> , the rows are features, and the columns are samples.
networkSets	A object of type <code>FeatureSetCollection</code> . The sets slot must contain a list of two-column matrices with each row corresponding to a binary interaction. Such sub-networks may be determined by a community detection algorithm.
absolute	If TRUE, then the absolute values of the differences are returned.
target	If measurements is a <code>MultiAssayExperiment</code> , the name of the data table to be used.
...	Variables not used by the <code>matrix</code> nor the <code>MultiAssayExperiment</code> method which are passed into and used by the <code>DataFrame</code> method.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

The pairs of features known to interact with each other are specified by `networkSets`.

**Value**

An object of class `DataFrame` with one column for each interactor pair difference and one row for each sample. Additionally, `mcols(resultTable)` provides a `DataFrame` with a column named "original" containing the name of the sub-network each meta-feature belongs to.

**Author(s)**

Dario Strbenac

**References**

Dynamic modularity in protein interaction networks predicts breast cancer outcome, Ian W Taylor, Rune Linding, David Warde-Farley, Yongmei Liu, Catia Pesquita, Daniel Faria, Shelley Bull, Tony Pawson, Quaid Morris and Jeffrey L Wrana, 2009, *Nature Biotechnology*, Volume 27 Issue 2, <https://www.nature.com/articles/nbt.1522>.

**Examples**

```
networksList <- list(`A Hub` = matrix(c('A', 'A', 'A', 'B', 'C', 'D'), ncol = 2),
                   `G Hub` = matrix(c('G', 'G', 'G', 'H', 'I', 'J'), ncol = 2))
netSets <- FeatureSetCollection(networksList)

# Differential correlation for sub-network with hub A.
measurements <- matrix(c(5.7, 10.1, 6.9, 7.7, 8.8, 9.1, 11.2, 6.4, 7.0, 5.5,
                        5.6, 9.6, 7.0, 8.4, 10.8, 12.2, 8.1, 5.7, 5.4, 12.1,
                        4.5, 9.0, 6.9, 7.0, 7.3, 6.9, 7.8, 7.9, 5.7, 8.7,
                        8.1, 10.6, 7.4, 7.15, 10.4, 6.1, 7.3, 2.7, 11.0, 9.1,
                        round(rnorm(60, 8, 1), 1)), ncol = 10, byrow = TRUE)
```

```
rownames(measurements) <- LETTERS[1:10]
colnames(measurements) <- paste("Patient", 1:10)

interactorDifferences(measurements, netSets)
```

---

kNNinterface

*An Interface for class Package's knn Function*


---

## Description

More details of k Nearest Neighbours are available in the documentation of [knn](#).

## Usage

```
## S4 method for signature 'matrix'
kNNinterface(measurements, classes, test, ...)
## S4 method for signature 'Dataframe'
kNNinterface(measurements, classes, test, ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
kNNinterface(measurements, test, targets = names(measurements), ...)
```

## Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">Dataframe</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples. If of type <a href="#">Dataframe</a> , the data set is subset to only those features of type <code>integer</code> .
classes	Either a vector of class labels of class <code>factor</code> of the same length as the number of samples in <code>measurements</code> or if the <code>measurements</code> are of class <code>Dataframe</code> a character vector of length 1 containing the column name in measurement is also permitted. Not used if <code>measurements</code> is a <code>MultiAssayExperiment</code> object.
test	An object of the same class as <code>measurements</code> with no samples in common with <code>measurements</code> and the same number of features as it.
targets	If <code>measurements</code> is a <code>MultiAssayExperiment</code> , the names of the data tables to be used. <code>"clinical"</code> is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the <code>matrix</code> nor the <code>MultiAssayExperiment</code> method which are passed into and used by the <code>Dataframe</code> method or parameters that <a href="#">knn</a> can accept.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

## Details

Data tables which consist entirely of non-numeric data cannot be analysed. If `measurements` is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `Dataframe` accessible by the `colData` function with column name `"class"`.



**Value**

A factor vector, the same as is returned by [knn](#).

**Author(s)**

Dario Strbenac

**Examples**

```
if(require(class))
{
  classes <- factor(rep(c("Healthy", "Disease"), each = 5),
                  levels = c("Healthy", "Disease"))
  measurements <- matrix(c(rnorm(50, 10), rnorm(50, 5)), ncol = 10)
  colnames(measurements) <- paste("Sample", 1:10)
  rownames(measurements) <- paste("mRNA", 1:10)
}

kNNinterface(measurements[, 1:9], classes[1:9], measurements[, 10, drop = FALSE])
```

---

KolmogorovSmirnovSelection

*Selection of Differential Distributions with Kolmogorov-Smirnov Distance*

---

**Description**

Ranks features by largest Kolmogorov-Smirnov distance and chooses the features which have best resubstitution performance.

**Usage**

```
## S4 method for signature 'matrix'
KolmogorovSmirnovSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
KolmogorovSmirnovSelection(measurements, classes, datasetName,
                           trainParams, predictParams, resubstituteParams, ...,
                           selectionName = "Kolmogorov-Smirnov Test", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
KolmogorovSmirnovSelection(measurements, targets = names(measurements), ...)
```

**Arguments**

**measurements** Either a [matrix](#), [DataFrame](#) or [MultiAssayExperiment](#) containing the training data. For a matrix, the rows are features, and the columns are samples.

**classes** Either a vector of class labels of class [factor](#) of the same length as the number of samples in `measurements` or if the measurements are of class `DataFrame` a character vector of length 1 containing the column name in measurement is also permitted. Not used if `measurements` is a `MultiAssayExperiment` object.

targets	If measurements is a MultiAssayExperiment, the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the matrix nor the MultiAssayExperiment method which are passed into and used by the DataFrame method or options which are accepted by the function <code>ks.test</code> .
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <code>TrainParams</code> describing the classifier to use for training.
predictParams	A container of class <code>PredictParams</code> describing how prediction is to be done.
resubstituteParams	An object of class <code>ResubstituteParams</code> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

Features are sorted in order of biggest distance to smallest. The top number of features is used in a classifier, to determine which number of features has the best resubstitution performance.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name "class".

### Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac

### Examples

```
# First 20 features have bimodal distribution for Poor class.
# Other 80 features have normal distribution for both classes.
set.seed(1984)
genesMatrix <- sapply(1:25, function(sample)
  {
    randomMeans <- sample(c(8, 12), 20, replace = TRUE)
    c(rnorm(20, randomMeans, 1), rnorm(80, 10, 1))
  }
)
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample) rnorm(100, 10, 1)))
rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
classes <- factor(rep(c("Poor", "Good"), each = 25))
```

```

resubstituteParams <- ResubstituteParams(nFeatures = seq(5, 25, 5),
                                       performanceType = "balanced error",
                                       better = "lower")
selected <- KolmogorovSmirnovSelection(genesMatrix, classes, "Example",
                                     trainParams = TrainParams(naiveBayesKernel),
                                     predictParams = PredictParams(NULL),
                                     resubstituteParams = resubstituteParams)

head(selected@chosenFeatures)
plotFeatureClasses(genesMatrix, classes, "Gene 13", dotBinWidth = 0.25,
                  xAxisLabel = bquote(log[2]*'(expression)'))

```

---

kTSPclassifier	<i>Classification Using k Pairs of Features With Relative Differences Between Classes</i>
----------------	---

---

## Description

Each pair of features votes for a class based on whether the value of one feature is less than the other feature.

## Usage

```

## S4 method for signature 'matrix'
kTSPclassifier(measurements, classes, test, featurePairs = NULL, ...)
## S4 method for signature 'DataFrame'
kTSPclassifier(measurements, classes, test, featurePairs = NULL,
              weighted = c("unweighted", "weighted", "both"),
              minDifference = 0, returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
kTSPclassifier(measurements, test, target = names(measurements)[1],
              featurePairs = NULL, ...)

```

## Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in <a href="#">measurements</a> or if the <a href="#">measurements</a> are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if <a href="#">measurements</a> is a <a href="#">MultiAssayExperiment</a> object.
test	An object of the same class as <a href="#">measurements</a> with no samples in common with <a href="#">measurements</a> and the same number of features as it.
featurePairs	An object of class as <a href="#">Pairs</a> containing the pairs of features to determine whether the inequality of the first feature being less than the second feature holds, indicating evidence for the second level of the classes factor.

target	If measurements is a MultiAssayExperiment, the name of the data table to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Unused variables by the methods for a matrix or a MultiAssayExperiment passed to the DataFrame method which does the classification.
weighted	Default: "both". Either "both", "unweighted" or "weighted". In weighted mode, the difference in densities is summed over all features. If unweighted mode, each feature's vote is worth the same. Both can be calculated simultaneously.
minDifference	Default: 0. The minimum difference in densities for a feature to be allowed to vote. Can be a vector of cutoffs. If no features for a particular sample have a difference large enough, the class predicted is simply the largest class.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, score for a sample belonging to the second class, as determined by the factor levels, or both labels and scores in a data.frame.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

If weighted is TRUE, then a sample's predicted class is based on the sum of differences of measurements for each feature pair. Otherwise, when weighted is FALSE, each pair of features has an equal vote, the predicted class is the one with the most votes. If the voting is tied, the the class with the most samples in the training set is voted for.

Because this method compares different features, they need to have comparable measurements. For example, RNA-seq counts would be unsuitable since these depend on the length of a feature, whereas F.P.K.M. values would be suitable.

The featurePairs to use is recommended to be determined in conjunction with pairsDifferencesSelection.

### Value

A vector or list of class prediction information, as long as the number of samples in the test data, or lists of such information, if a variety of predictions is generated.

### Author(s)

Dario Strbenac

### See Also

[pairsDifferencesSelection](#) for a function which could be used to do feature selection before the k-TSP classifier is run.

**Examples**

```
# Difference in differences for features A and C between classes.
measurements <- matrix(c(9.9, 10.5, 10.1, 10.9, 11.0, 6.6, 7.7, 7.0, 8.1, 6.5,
                        8.5, 10.5, 12.5, 10.5, 9.5, 8.5, 10.5, 12.5, 10.5, 9.5,
                        6.6, 7.7, 7.0, 8.1, 6.5, 11.2, 11.0, 11.1, 11.4, 12.0,
                        8.1, 10.6, 7.4, 7.1, 10.4, 6.1, 7.3, 2.7, 11.0, 9.1,
                        round(rnorm(60, 8, 1), 1)), ncol = 10, byrow = TRUE)
classes <- factor(rep(c("Good", "Poor"), each = 5))

rownames(measurements) <- LETTERS[1:10]
colnames(measurements) <- names(classes) <- paste("Patient", 1:10)

trainIndex <- c(1:4, 6:9)
trainMatrix <- measurements[, trainIndex]
testMatrix <- measurements[, c(5, 10)]

featurePairs <- Pairs('A', 'C') # Could be selected by pairsDifferencesSelection function.
kTSPclassifier(trainMatrix, classes[trainIndex], testMatrix, featurePairs)
```

---

**KullbackLeiblerSelection***Selection of Differential Distributions with Kullback-Leibler Distance*

---

**Description**

Ranks features by largest Kullback-Leibler distance and chooses the features which have best re-substitution performance.

**Usage**

```
## S4 method for signature 'matrix'
KullbackLeiblerSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
KullbackLeiblerSelection(measurements, classes, datasetName,
                        trainParams, predictParams, resubstituteParams, ...,
                        selectionName = "Kullback-Leibler Divergence", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
KullbackLeiblerSelection(measurements, targets = names(measurements), ...)
```

**Arguments**

**measurements** Either a [matrix](#), [DataFrame](#) or [MultiAssayExperiment](#) containing the training data. For a matrix, the rows are features, and the columns are samples.

**classes** Either a vector of class labels of class [factor](#) of the same length as the number of samples in `measurements` or if the measurements are of class `DataFrame` a character vector of length 1 containing the column name in measurement is also permitted. Not used if `measurements` is a `MultiAssayExperiment` object.

targets	If measurements is a MultiAssayExperiment, the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the matrix nor the MultiAssayExperiment method which are passed into and used by the DataFrame method or options which are accepted by the function <a href="#">getLocationsAndScales</a> .
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

The distance is defined as  $\frac{1}{2} \times \left( \frac{(location_1 - location_2)^2}{scale_1^2} + \frac{(location_1 - location_2)^2}{scale_2^2} + \frac{scale_2^2}{scale_1^2} + \frac{scale_1^2}{scale_2^2} \right)$

The subscripts denote the group which the parameter is calculated for.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class MultiAssayExperiment, the factor of sample classes must be stored in the DataFrame accessible by the colData function with column name "class".

### Value

An object of class [SelectResult](#) or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac

### Examples

```
# First 20 features have bimodal distribution for Poor class.
# Other 80 features have normal distribution for both classes.
genesMatrix <- sapply(1:25, function(sample)
  {
    randomMeans <- sample(c(8, 12), 20, replace = TRUE)
    c(rnorm(20, randomMeans, 1), rnorm(80, 10, 1))
  }
)
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample) rnorm(100, 10, 1)))
classes <- factor(rep(c("Poor", "Good"), each = 25))

resubstituteParams <- ResubstituteParams(nFeatures = seq(5, 25, 5),
  performanceType = "balanced error",
```

```

                                better = "lower")
KullbackLeiblerSelection(genesMatrix, classes, "Example",
                        trainParams = TrainParams(naiveBayesKernel),
                        predictParams = PredictParams(NULL),
                        resubstituteParams = resubstituteParams
                        )

```

leveneSelection

*Selection of Differential Variability with Levene Statistic***Description**

Ranks features by largest Levene statistic and chooses the features which have best resubstitution performance.

**Usage**

```

## S4 method for signature 'matrix'
leveneSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
leveneSelection(measurements, classes, datasetName,
                trainParams, predictParams, resubstituteParams,
                selectionName = "Levene Test", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
leveneSelection(measurements, targets = names(measurements), ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the matrix nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.

**selectionName** A name to identify this selection method by. Stored in the result.

**verbose** Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

Levene's statistic for unequal variance between groups is a robust version of Bartlett's statistic.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name "class".

### Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac

### Examples

```
# First 20 features have bimodal distribution for Poor class.
# Other 80 features have normal distribution for both classes.
set.seed(1984)
genesMatrix <- sapply(1:25, function(sample)
  {
    randomMeans <- sample(c(8, 12), 20, replace = TRUE)
    c(rnorm(20, randomMeans, 1), rnorm(80, 10, 1))
  }
)
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample) rnorm(100, 10, 1)))
rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
classes <- factor(rep(c("Poor", "Good"), each = 25))
genesMatrix <- subtractFromLocation(genesMatrix, 1:ncol(genesMatrix))

resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
  performanceType = "balanced error",
  better = "lower")
selected <- leveneSelection(genesMatrix, classes, "Example",
  trainParams = TrainParams(fisherDiscriminant),
  predictParams = PredictParams(NULL),
  resubstituteParams = resubstituteParams)

selected@chosenFeatures
```



---

likelihoodRatioSelection

*Selection of Differential Distributions with Likelihood Ratio Statistic*


---

## Description

Ranks features by largest ratio and chooses the features which have the best resubstitution performance.

## Usage

```
## S4 method for signature 'matrix'
likelihoodRatioSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
likelihoodRatioSelection(measurements, classes, datasetName,
                        trainParams, predictParams, resubstituteParams,
                        alternative = c(location = "different", scale = "different"),
                        ..., selectionName = "Likelihood Ratio Test (Normal)", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
likelihoodRatioSelection(measurements, targets = names(measurements), ...)
```

## Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or options which are accepted by the function <a href="#">getLocationsAndScales</a> .
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
alternative	Default: <code>c("different", "different")</code> . A vector of length 2. The first element specifies the location of the alternate hypothesis. The second element specifies the scale of the alternate hypothesis. Valid values in each element are "same" or "different".

`selectionName` A name to identify this selection method by. Stored in the result.

`verbose` Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

## Details

Likelihood ratio test of null hypothesis that the location and scale are the same for both groups, and an alternate hypothesis that is specified by parameters. The location and scale of features is calculated by `getLocationsAndScales`. The distribution fitted to the data is the normal distribution.

Data tables which consist entirely of non-numeric data cannot be analysed. If measurements is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name "class".

## Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

## Author(s)

Dario Strbenac

## Examples

```
# First 20 features have bimodal distribution for Poor class.
# Other 80 features have normal distribution for both classes.
set.seed(1984)
genesMatrix <- sapply(1:25, function(sample)
  {
    randomMeans <- sample(c(8, 12), 20, replace = TRUE)
    c(rnorm(20, randomMeans, 1), rnorm(80, 10, 1))
  }
)
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample) rnorm(100, 10, 1)))
rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
classes <- factor(rep(c("Poor", "Good"), each = 25))

resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
  performanceType = "balanced error",
  better = "lower")
selected <- likelihoodRatioSelection(genesMatrix, classes, "Example",
  trainParams = TrainParams(naiveBayesKernel),
  predictParams = PredictParams(NULL),
  resubstituteParams = resubstituteParams)

head(selected@chosenFeatures[[1]])
```

---

 limmaSelection      *Selection of Differentially Abundant Features*


---

**Description**

Uses a moderated t-test with empirical Bayes shrinkage to select differentially expressed features.

**Usage**

```
## S4 method for signature 'matrix'
limmaSelection(measurements, classes, ...)
## S4 method for signature 'DataFrame'
limmaSelection(measurements, classes, datasetName,
               trainParams, predictParams, resubstituteParams, ...,
               selectionName = "Moderated t-test", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
limmaSelection(measurements, targets = NULL, ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	A vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	Names of data tables to be combined into a single table and used in the analysis.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or optional settings that are passed to <a href="#">lmFit</a> .
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.
resubstituteParams	An object of class <a href="#">ResubstituteParams</a> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

This selection method looks for changes in means and uses a moderated t-test.

**Value**

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

**Author(s)**

Dario Strbenac

**References**

Limma: linear models for microarray data, Gordon Smyth, 2005, In: Bioinformatics and Computational Biology Solutions using R and Bioconductor, Springer, New York, pages 397-420.

**Examples**

```
#if(require(sparsediscrim))
#{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))

  resubstituteParams <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "balanced error", better = "lower")
  selected <- limmaSelection(genesMatrix, classes, "Example",
    trainParams = TrainParams(), predictParams = PredictParams(),
    resubstituteParams = resubstituteParams)

  selected@chosenFeatures
#}
```

---

listOrCharacterOrNULL *Union of a List and a Character Vector and NULL*

---

**Description**

Allows a slot to be either a list, character or a NULL.

**Author(s)**

Dario Strbenac

**Examples**

```
setClass("HardClassifier", representation(model = "listOrCharacterOrNULL"))
classifier <- new("HardClassifier", model = "Good") # Optimistic classifier.
```

---

listOrNULL

*Union of a List and NULL*


---

**Description**

Allows a slot to be either a list or a NULL.

**Author(s)**

Dario Strbenac

**Examples**

```
setClass("EasyClassifier", representation(model = "listOrNULL"))
classifier <- new("EasyClassifier", model = NULL) # Optimistic classifier.
```

---

mixmodels

*Classification based on Differential Distribution utilising Mixtures of Normals*


---

**Description**

Fits mixtures of normals for every feature, separately for each class.

**Usage**

```
## S4 method for signature 'matrix'
mixModelsTrain(measurements, ...)
## S4 method for signature 'DataFrame'
mixModelsTrain(measurements, classes, ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
mixModelsTrain(measurements, targets = names(measurements), ...)
## S4 method for signature 'MixModelsListsSet,matrix'
mixModelsPredict(models, test, ...)
## S4 method for signature 'MixModelsListsSet,DataFrame'
mixModelsPredict(models, test, weighted = c("unweighted", "weighted", "both"),
  weight = c("height difference", "crossover distance", "both"),
  densityXvalues = 1024, minDifference = 0,
  returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'MixModelsListsSet,MultiAssayExperiment'
mixModelsPredict(models, test, targets = names(test), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in <a href="#">measurements</a> or if the <a href="#">measurements</a> are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in <a href="#">measurement</a> is also permitted. Not used if <a href="#">measurements</a> is a <a href="#">MultiAssayExperiment</a> object.
test	An object of the same class as <a href="#">measurements</a> with no samples in common with <a href="#">measurements</a> and the same number of features as it. Also, if a <a href="#">DataFrame</a> , the <a href="#">class</a> column must be absent.
targets	If <a href="#">measurements</a> is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or extra arguments for training passed to <a href="#">mixmodCluster</a> . The argument <a href="#">nbCluster</a> is mandatory.
models	A <a href="#">MixModelsListsSet</a> of models generated by the training function and training class information. There is one element for each class. Another element at the end of the list has the class sizes of the classes in the training data.
weighted	Default: "unweighted". Either "unweighted", "weighted" or "both". In weighted mode, the difference in densities is summed over all features. If unweighted mode, each feature's vote is worth the same. Both can be calculated simultaneously.
weight	Default: "both". Either "both", "height difference", or "crossover distance". The type of weight to calculate. For "height difference", the weight of each prediction is equal to the sum of the vertical distances for all of the mixture components within one class subtracted from the sum of the components of the other class, summed for each value of x. For "crossover distance", the x positions where the mixture density of the class being considered crosses another class' density is firstly calculated. The predicted class is the class with the highest mixture sum at the particular value of x and the weight is the distance of x from the nearest density crossover point.
densityXvalues	Default: 1024. Only relevant when <a href="#">weight</a> is "crossover distance". The number of equally-spaced locations at which to calculate y values for each mixture density.
minDifference	Default: 0. The minimum difference in sums of mixture densities between the class with the highest sum and the class with the second highest sum for a feature to be allowed to vote. Can be a vector of cutoffs. If no features for a particular sample have a difference large enough, the class predicted is simply the largest class.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of predicted classes, a matrix of scores with columns corresponding to classes, as determined by the factor levels of classes, or both a column of predicted classes and columns of class scores in a <code>data.frame</code> .

`verbose` Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

If `weighted` is TRUE, then a sample's predicted class is the class with the largest sum of weights, each scaled for the number of samples in the training data of each class. Otherwise, when `weighted` is FALSE, each feature has an equal vote, and votes for the class with the largest weight, scaled for class sizes in the training set.

If `weight` is "crossover distance", the crossover points are computed by considering the distance between `y` values of the two densities at every `x` value. `x` values for which the sign of the difference changes compared to the difference of the closest lower value of `x` are used as the crossover points.

### Value

For `mixModelsTrain`, a list of trained models of class `MixmodCluster`. For `mixModelsPredict`, a vector or list of class prediction information (i.e. classes and/or scores), as long as the number of samples in the test data, or lists of such information, if both `weighted` and `unweighted` voting was used or a range of `minDifference` values was provided.

### Author(s)

Dario Strbenac

### Examples

```
# First 25 samples and first 5 genes are mixtures of two normals. Last 25 samples are
# one normal.

genesMatrix <- sapply(1:25, function(geneColumn) c(rnorm(5, sample(c(5, 15), replace = TRUE, 5))))
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(geneColumn) c(rnorm(5, 9, 1))))
genesMatrix <- rbind(genesMatrix, sapply(1:50, function(geneColumn) rnorm(5, 9, 1)))
rownames(genesMatrix) <- paste("Gene", 1:10)
colnames(genesMatrix) <- paste("Sample", 1:50)
classes <- factor(rep(c("Poor", "Good"), each = 25), levels = c("Good", "Poor"))

trainSamples <- c(1:15, 26:40)
testSamples <- c(16:25, 41:50)
selected <- 1:5

trained <- mixModelsTrain(genesMatrix[selected, trainSamples], classes[trainSamples],
                        nbCluster = 1:3)
mixModelsPredict(trained, genesMatrix[selected, testSamples], minDifference = 0:3)
```

---

MixModelsListsSet      *Container for a List of Lists Containing Mixture Models*

---

**Description**

Stores a list of lists of trained mixture models, to prevent them being unintentionally being unlisted during cross-validation. Not intended for end-user.

**Constructor**

MixModelsListsSet(set)

Creates a MixModelsListsSet object which stores the mixture models.

set A list as long as the number of classes in the data set. Each element is a list, which each element of is a mixture model trained on one feature.

**Author(s)**

Dario Strbenac

**Examples**

```
if(require(Rmixmod))
{
  mixModels <- list(Good = list(mixmodCluster(rnorm(20), nbCluster = 1:2)),
                  Poor = list(mixmodCluster(rnorm(20), nbCluster = 1:2)))
  MixModelsListsSet(mixModels)
}
```

---

multnet      *Trained multnet Object*

---

**Description**

Enables S4 method dispatching on it.

**Author(s)**

Dario Strbenac



---

naiveBayesKernel	<i>Classification Using A Bayes Classifier with Kernel Density Estimates</i>
------------------	--

---

## Description

Kernel density estimates are fitted to the training data and a naive Bayes classifier is used to classify samples in the test data.

## Usage

```
## S4 method for signature 'matrix'
naiveBayesKernel(measurements, classes, test, ...)
## S4 method for signature 'DataFrame'
naiveBayesKernel(measurements, classes, test,
  densityFunction = density,
  densityParameters = list(bw = "nrd0", n = 1024,
    from = expression(min(featureValues)),
    to = expression(max(featureValues))),
  weighted = c("unweighted", "weighted", "both"),
  weight = c("height difference", "crossover distance", "both"),
  minDifference = 0, returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
naiveBayesKernel(measurements, test, targets = names(measurements), ...)
```

## Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Unused variables by the three top-level methods passed to the internal method which does the classification.
densityFunction	Default: <a href="#">density</a> . A function which will return a probability density, which is essentially a list with x and y coordinates.
densityParameters	A list of options for densityFunction. Default: <code>list(bw = "nrd0", n = 1024, from = expression(min(featureValues)), to = expression(max(featureValues)))</code> .

weighted	Default: "unweighted". Either "unweighted", "weighted" or "both". In weighted mode, the difference in densities is summed over all features. If unweighted mode, each feature's vote is worth the same. Both can be calculated simultaneously.
weight	Default: "both". Either "both", "height difference", or "crossover distance". The type of weight to calculate. For "height difference", the weight of each prediction is equal to the vertical distance between the highest density and the second-highest, for a particular value of x. For "crossover distance", the x positions where two densities cross is firstly calculated. The predicted class is the class with the highest density at the particular value of x and the weight is the distance of x from the nearest density crossover point.
minDifference	Default: 0. The minimum difference in density height between the highest density and second-highest for a feature to be allowed to vote. Can be a vector of cutoffs. If no features for a particular sample have a difference large enough, the class predicted is simply the largest class.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of predicted classes, a matrix of scores with columns corresponding to classes, as determined by the factor levels of classes, or both a column of predicted classes and columns of class scores in a <code>data.frame</code> .
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

If `weighted` is `TRUE`, then a sample's predicted class is the class with the largest sum of weights, each scaled for the number of samples in the training data of each class. Otherwise, when `weighted` is `FALSE`, each feature has an equal vote, and votes for the class with the largest weight, scaled for class sizes in the training set.

The variable name of each feature's measurements in the iteration over all features is `featureValues`. This is important to know if each feature's measurements need to be referred to in the specification of `densityParameters`, such as for specifying the range of x values of the density function to be computed. For example, see the default value of `densityParameters` above.

If `weight` is "crossover distance", the crossover points are computed by considering the distance between y values of all of the densities at every x value. x values for which a class density crosses any other class' density are used as the crossover points for that class.

### Value

A vector or list of class prediction information (i.e. classes and/or scores), as long as the number of samples in the test data, or lists of such information, if both weighted and unweighted voting was used or a range of `minDifference` values was provided.

### Author(s)

Dario Strbenac, John Ormerod

**Examples**

```

trainMatrix <- matrix(rnorm(1000, 8, 2), ncol = 10)
classes <- factor(rep(c("Poor", "Good"), each = 5))

# Make first 30 genes increased in value for poor samples.
trainMatrix[1:30, 1:5] <- trainMatrix[1:30, 1:5] + 5

testMatrix <- matrix(rnorm(1000, 8, 2), ncol = 10)

# Make first 30 genes increased in value for sixth to tenth samples.
testMatrix[1:30, 6:10] <- testMatrix[1:30, 6:10] + 5

naiveBayesKernel(trainMatrix, classes, testMatrix)

```

---

networkCorrelationsSelection

*Selection of Differentially Correlated Hub Sub-networks*


---

**Description**

Ranks sub-networks by largest within-class to between-class correlation variability and chooses the sub-networks which have the best resubstitution performance.

**Usage**

```

## S4 method for signature 'matrix'
networkCorrelationsSelection(measurements, classes, metaFeatures = NULL, ...)
## S4 method for signature 'DataFrame'
networkCorrelationsSelection(measurements, classes, metaFeatures = NULL,
                             featureSets, datasetName, trainParams, predictParams, resubstituteParams,
                             selectionName = "Differential Correlation of Sub-networks", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
networkCorrelationsSelection(measurements, target = NULL, metaFeatures = NULL, ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
metaFeatures	A <a href="#">DataFrame</a> with the same number of samples as the numeric table of interest. The number of derived features in this table will be different to the original input data table. The command <code>mcols(metaFeatures)</code> must return a <a href="#">DataFrame</a> which has an "original" column with as many rows as there are meta-features and specifies the feature which the meta-feature is originally derived from (e.g. network name).

featureSets	A object of type <code>FeatureSetCollection</code> . The sets slot must contain a list of two-column matrices with each row corresponding to a binary interaction. Such sub-networks may be determined by a community detection algorithm. This will be used to determine which features belong to which sub-networks before calculating a statistic for each sub-network.
target	If measurements is a <code>MultiAssayExperiment</code> , the name of the data table to be used.
...	Variables not used by the <code>matrix</code> nor the <code>MultiAssayExperiment</code> method which are passed into and used by the <code>DataFrame</code> method.
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <code>TrainParams</code> describing the classifier to use for training.
predictParams	A container of class <code>PredictParams</code> describing how prediction is to be done.
resubstituteParams	An object of class <code>ResubstituteParams</code> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

## Details

The selection of sub-networks is based on the average difference in correlations between each pair of interactors, considering the samples within each class separately. Such differences of correlations within each of the two classes are scaled by the average difference of correlations within each class.

More formally, let  $C_{i,j}$  be the correlation of the  $j^{\text{th}}$  edge using all samples belonging to class  $i$ . Then, let  $\bar{C}_{i\bullet}$  be defined as  $\frac{C_{i,1}+C_{i,2}+\dots+C_{i,e}}{n}$  where  $e$  is the number of edges in the sub-network being considered. Also, let  $\bar{C}_{\bullet\bullet}$ , the average overall correlation, be  $\frac{C_{1\bullet}+C_{2\bullet}}{2}$ . Then, the between-class sum-of-squares (BSS) is  $\sum_{i=1}^2 e(\bar{C}_{i\bullet} - \bar{C}_{\bullet\bullet})^2$ . Also the within-class sum-of-squares (WSS) is  $\sum_{i=1}^2 \sum_{j=1}^e (C_{i,j} - \bar{C}_{i\bullet})^2$ . The sub-networks are ranked in decreasing order of  $\frac{BSS}{WSS}$ .

The classifier specified by `trainParams` and `predictParams` is used to calculate resubstitution error rates using the transformation of the data set provided by `metaFeatures`. The set of top-ranked sub-networks which give the lowest resubstitution error rate are finally selected.

Data tables which consist entirely of non-numeric data cannot be analysed. If `measurements` is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name "class".

## Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

## Author(s)

Dario Strbenac

## References

Network-based biomarkers enhance classical approaches to prognostic gene expression signatures, Rebecca L Barter, Sarah-Jane Schramm, Graham J Mann and Yee Hwa Yang, 2014, *BMC Systems Biology*, Volume 8 Supplement 4 Article S5, <https://bmcsystbiol.biomedcentral.com/articles/10.1186/1752-0509-8-S4-S5>.

## See Also

[interactorDifferences](#) for an example of a function which can turn the measurements into meta-features for classification.

## Examples

```
networksList <- list(`A Hub` = matrix(c('A', 'A', 'A', 'B', 'C', 'D'), ncol = 2),
  `G Hub` = matrix(c('G', 'G', 'G', 'H', 'I', 'J'), ncol = 2))
netSets <- FeatureSetCollection(networksList)

# Differential correlation for sub-network with hub A.
measurements <- matrix(c(5.7, 10.1, 6.9, 7.7, 8.8, 9.1, 11.2, 6.4, 7.0, 5.5,
  5.6, 9.6, 7.0, 8.4, 10.8, 12.2, 8.1, 5.7, 5.4, 12.1,
  4.5, 9.0, 6.9, 7.0, 7.3, 6.9, 7.8, 7.9, 5.7, 8.7,
  8.1, 10.6, 7.4, 7.1, 10.4, 6.1, 7.3, 2.7, 11.0, 9.1,
  round(rnorm(60, 8, 1), 1)), ncol = 10, byrow = TRUE)
classes <- factor(rep(c("Good", "Poor"), each = 5))

rownames(measurements) <- LETTERS[1:10]
colnames(measurements) <- names(classes) <- paste("Patient", 1:10)

Idifferences <- interactorDifferences(measurements, netSets)

# The features are sub-networks and there are only two in this example.
resubstituteParams <- ResubstituteParams(nFeatures = 1:2,
  performanceType = "balanced error", better = "lower")

predictParams <- PredictParams(NULL)
networkCorrelationsSelection(measurements, classes, metaFeatures = Idifferences,
  featureSets = netSets, datasetName = "Example",
  trainParams = TrainParams(naiveBayesKernel),
  predictParams = predictParams,
  resubstituteParams = resubstituteParams)
```

---

NSCpredictInterface     *Interface for pamr.predict Function from pamr CRAN Package*

---

## Description

Restructures variables from ClassifyR framework to be compatible with [pamr.predict](#) definition.

**Usage**

```

## S4 method for signature 'pamrtrained,matrix'
NSCpredictInterface(trained, test, ...)
## S4 method for signature 'pamrtrained,DataFrame'
NSCpredictInterface(trained, test, classes = NULL, ..., returnType = c("both", "class", "score"), verbose = FALSE)
## S4 method for signature 'pamrtrained,MultiAssayExperiment'
NSCpredictInterface(trained, test, targets = names(test), ...)

```

**Arguments**

<code>trained</code>	An object of class <code>pamrtrained</code> .
<code>test</code>	An object of the same class as <code>measurements</code> with no samples in common with <code>measurements</code> used in the training stage and the same number of features as it. Also, if a <code>DataFrame</code> , the <code>class</code> column must be absent.
<code>classes</code>	Either <code>NULL</code> or a character vector of length 1, specifying the column name to remove.
<code>targets</code>	If <code>test</code> is a <code>MultiAssayExperiment</code> , the names of the data tables to be used. <code>"clinical"</code> is also a valid value and specifies that numeric variables from the clinical data table will be used.
<code>...</code>	Variables not used by the <code>matrix</code> nor the <code>MultiAssayExperiment</code> method which are passed into and used by the <code>DataFrame</code> method or optional settings that are passed to <code>pamr.predict</code> .
<code>returnType</code>	Default: <code>"both"</code> . Either <code>"class"</code> , <code>"score"</code> or <code>"both"</code> . Sets the return value from the prediction to either a vector of class labels, score for a sample belonging to the second class, as determined by the factor levels, or both labels and scores in a <code>data.frame</code> .
<code>verbose</code>	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

This function is an interface between the `ClassifyR` framework and `pamr.predict`. It selects the highest threshold that gives the minimum error rate in the training data.

**Value**

Either a factor vector of predicted classes, a matrix of scores for each class, or a table of both the class labels and class scores, depending on the setting of `returnType`.

**Author(s)**

Dario Strbenac

**See Also**

`pamr.predict` for the function that was interfaced to.

**Examples**

```

if(require(pamr))
{
  # Samples in one class with differential expression to other class.
  genesMatrix <- sapply(1:25, function(geneColumn) c(rnorm(100, 9, 1)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(geneColumn)
    c(rnorm(75, 9, 1), rnorm(25, 14, 1))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))

  fit <- NSCtrainInterface(genesMatrix[, c(1:20, 26:45)], classes[c(1:20, 26:45)])
  NSCpredictInterface(fit, genesMatrix[, c(21:25, 46:50)])
}

```

---

NSCselectionInterface *Interface for pamr.listgenes Function from pamr CRAN Package*

---

**Description**

Restructures variables from ClassifyR framework to be compatible with [pamr.listgenes](#) definition.

**Usage**

```

## S4 method for signature 'matrix'
NSCselectionInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
NSCselectionInterface(measurements, classes, datasetName,
  trained, ..., selectionName = "Shrunken Centroids", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
NSCselectionInterface(measurements, targets = names(measurements), ...)

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
datasetName	A name for the data set used. Stored in the result.
trained	The output of <a href="#">NSCtrainInterface</a> , which is identical to the output of <a href="#">pamr.listgenes</a> .
...	Variables not used by the matrix nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or extra arguments passed to <a href="#">pamr.listgenes</a> .

selectionName A name to identify this selection method by. Stored in the result.

verbose Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

This function is an interface between the ClassifyR framework and [pamr.listgenes](#).

The set of features chosen is the obtained by considering the range of thresholds provided to [NSCtrainInterface](#) and using the threshold that obtains the lowest cross-validation error rate on the training set.

### Value

An object of class [SelectResult](#). The rankedFeatures slot will be empty.

### Author(s)

Dario Strbenac

### See Also

[pamr.listgenes](#) for the function that was interfaced to.

### Examples

```
if(require(pamr))
{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(geneColumn) c(rnorm(100, 9, 1)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(geneColumn)
    c(rnorm(75, 9, 1), rnorm(25, 14, 1))))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
  classes <- factor(rep(c("Poor", "Good"), each = 25))

  trained <- NSCtrainInterface(genesMatrix, classes)
  selected <- NSCselectionInterface(genesMatrix, classes, "Example", trained)

  selected@chosenFeatures
}
```

---

NSCtrainInterface      *Interface for pamr.train Function from pamr CRAN Package*

---

### Description

Restructures variables from ClassifyR framework to be compatible with [pamr.train](#) definition.



**Usage**

```
## S4 method for signature 'matrix'
NSCtrainInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
NSCtrainInterface(measurements, classes, ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
NSCtrainInterface(measurements, targets = names(measurements), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method or extra arguments passed to <a href="#">pamr.train</a> .
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Details**

This function is an interface between the [ClassifyR](#) framework and [pamr.train](#).

**Value**

A list with elements as described in [pamr.train](#).

**Author(s)**

Dario Strbenac

**See Also**

[pamr.train](#) for the function that was interfaced to.

**Examples**

```
if(require(pamr))
{
  # Samples in one class with differential expression to other class.
  genesMatrix <- sapply(1:25, function(geneColumn) c(rnorm(100, 9, 1)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(geneColumn)
```

```

                                c(rnorm(75, 9, 1), rnorm(25, 14, 1)))
classes <- factor(rep(c("Poor", "Good"), each = 25))

NSCtrainInterface(genesMatrix, classes)
}

```

---

pairsDifferencesSelection

*Selection of Pairs of Features that are Different Between Classes*

---

### Description

Ranks pre-specified pairs of features by the largest difference of the sum of measurement differences over all samples within a class and chooses the pairs of features which have the best resubstitution performance.

### Usage

```

## S4 method for signature 'matrix'
pairsDifferencesSelection(measurements, classes, featurePairs = NULL, ...)
## S4 method for signature 'DataFrame'
pairsDifferencesSelection(measurements, classes, featurePairs = NULL,
                          datasetName, trainParams, predictParams, resubstituteParams,
                          selectionName = "Pairs Differences", verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
pairsDifferencesSelection(measurements, target = names(measurements)[1], featurePairs = NULL, ...)

```

### Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in <a href="#">measurements</a> or if the <a href="#">measurements</a> are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in <a href="#">measurement</a> is also permitted. Not used if <a href="#">measurements</a> is a <a href="#">MultiAssayExperiment</a> object.
featurePairs	An S4 object of type <a href="#">Pairs</a> containing feature identifiers to calculate the sum of differences within each class for.
target	If <a href="#">measurements</a> is a <a href="#">MultiAssayExperiment</a> , the name of the data table to be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
datasetName	A name for the data set used. Stored in the result.
trainParams	A container of class <a href="#">TrainParams</a> describing the classifier to use for training.
predictParams	A container of class <a href="#">PredictParams</a> describing how prediction is to be done.

resubstituteParams	An object of class <code>ResubstituteParams</code> describing the performance measure to consider and the numbers of top features to try for resubstitution classification.
selectionName	A name to identify this selection method by. Stored in the result.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

Instead of considering whether one feature in a pair of features is consistently lower or higher than the other in the pair, this method takes the sum of differences across all samples within a class, to prevent ties in the ranking of pairs of features.

### Value

An object of class `SelectResult` or a list of such objects, if the classifier which was used for determining the specified performance metric made a number of prediction varieties.

### Author(s)

Dario Strbenac

### References

Simple decision rules for classifying human cancers from gene expression profiles, Aik C Tan, Daniel Q Naiman, Lei Xu, Raimond L. Winslow and Donald Geman, 2005, *Bioinformatics*, Volume 21 Issue 20, <https://academic.oup.com/bioinformatics/article/21/20/3896/203010>.

### See Also

`KTSPclassifier` for a classifier which makes use of the pairs of selected features in classification.

### Examples

```
featurePairs <- Pairs(c('A', 'A'), c('B', 'C'))

# Difference in differences for features A and C between classes.
measurements <- matrix(c(9.9, 10.5, 10.1, 10.9, 11.0, 6.6, 7.7, 7.0, 8.1, 6.5,
                        8.5, 10.5, 12.5, 10.5, 9.5, 8.5, 10.5, 12.5, 10.5, 9.5,
                        6.6, 7.7, 7.0, 8.1, 6.5, 11.2, 11.0, 11.1, 11.4, 12.0,
                        8.1, 10.6, 7.4, 7.1, 10.4, 6.1, 7.3, 2.7, 11.0, 9.1,
                        round(rnorm(60, 8, 1), 1)), ncol = 10, byrow = TRUE)
classes <- factor(rep(c("Good", "Poor"), each = 5))

rownames(measurements) <- LETTERS[1:10]
colnames(measurements) <- names(classes) <- paste("Patient", 1:10)

# The features are pairs and there are only two in this example.
resubstituteParams <- ResubstituteParams(nFeatures = 1:2,
                                         performanceType = "balanced error", better = "lower")
```

```

predictParams <- PredictParams(NULL)
pairsDifferencesSelection(measurements, classes, featurePairs = featurePairs,
                          datasetName = "Example",
                          trainParams = TrainParams(kTSPclassifier),
                          predictParams = predictParams,
                          resubstituteParams = resubstituteParams)

```

---

pamrtrained

*Trained pamr Object*

---

### Description

Enables S4 method dispatching on it.

### Author(s)

Dario Strbenac

---

performancePlot

*Plot Performance Measures for Various Classifications*

---

### Description

Draws a graphical summary of a particular performance measure for a list of classifications

### Usage

```

## S4 method for signature 'list'
performancePlot(results, aggregate = character(),
                xVariable = c("classificationName", "datasetName", "selectionName",
                              "validation"),
                performanceName = NULL,
                boxFillColouring = c("classificationName", "datasetName", "selectionName",
                                      "validation", "None"),
                boxFillColours = NULL,
                boxLineColouring = c("classificationName", "datasetName", "selectionName",
                                      "validation", "None"),
                boxLineColours = NULL,
                rowVariable = c("None", "validation", "datasetName", "classificationName",
                               "selectionName"),
                columnVariable = c("datasetName", "classificationName", "validation",
                                   "selectionName", "None"),
                yLimits = c(0, 1), fontSizes = c(24, 16, 12, 12), title = NULL,
                xLabel = "Analysis", yLabel = performanceName,
                margin = grid::unit(c(0, 0, 0, 0), "lines"), rotate90 = FALSE,
                showLegend = TRUE, plot = TRUE)

```

**Arguments**

results	A list of <code>ClassifyResult</code> objects.
aggregate	A character vector of the levels of <code>xVariable</code> to aggregate to a single number by taking the mean. This is particularly meaningful when the cross-validation is leave-k-out, when k is small.
xVariable	The factor to make separate boxes for.
performanceName	The name of the performance measure to make comparisons of. This is one of the names printed in the Performance Measures field when a <code>ClassifyResult</code> object is printed.
boxFillColouring	A factor to colour the boxes by.
boxFillColours	A vector of colours, one for each level of <code>boxFillColouring</code> .
boxLineColouring	A factor to colour the box lines by.
boxLineColours	A vector of colours, one for each level of <code>boxLineColouring</code> .
rowVariable	The slot name that different levels of are plotted as separate rows of boxplots.
columnVariable	The slot name that different levels of are plotted as separate columns of boxplots.
yLimits	The minimum and maximum value of the performance metric to plot.
fontSizes	A vector of length 4. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values. The fourth number is the font size of the titles of grouped plots, if any are produced. In other words, when <code>rowVariable</code> or <code>columnVariable</code> are not NULL.
title	An overall title for the plot.
xLabel	Label to be used for the x-axis.
yLabel	Label to be used for the y-axis of overlap percentages.
margin	The margin to have around the plot.
rotate90	Logical. IF TRUE, the plot is horizontal.
showLegend	If TRUE, a legend is plotted next to the plot. If FALSE, it is hidden.
plot	Logical. IF TRUE, a plot is produced on the current graphics device.

**Details**

Possible values for slot names are "datasetName", "classificationName", and "validation". If "None", then that graphic element is not used.

If there are multiple values for a performance measure in a single result object, it is plotted as a boxplot, unless `aggregate` is TRUE, in which case the all predictions in a single result object are considered simultaneously, so that only one performance number is calculated, and a barchart is plotted.

**Value**

An object of class `ggplot` and a plot on the current graphics device, if `plot` is TRUE.

**Author(s)**

Dario Strbenac

**Examples**

```

predicted <- list(data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))))
actual <- factor(rep(c("Healthy", "Cancer"), each = 5))
result1 <- ClassifyResult("Example", "Differential Expression", "t-test",
                        LETTERS[1:10], LETTERS[10:1], 100, list(1:100, c(1:9, 11:101)),
                        list(c(1:3), c(2, 5, 6), c(1:4), c(5:8), 1:5),
                        list(function(oracle){}), predicted, actual,
                        validation = list("permuteFold", 2, 2))
result1 <- calcCVperformance(result1, "macro F1")

predicted <- list(data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))),
                 data.frame(sample = sample(LETTERS[1:10], 20, replace = TRUE),
                           class = factor(rep(c("Healthy", "Cancer"), each = 10))))
result2 <- ClassifyResult("Example", "Differential Variability", "Bartlett Test",
                        LETTERS[1:10], LETTERS[10:1], 100, list(1:100, c(1:5, 11:105)),
                        list(c(1:3), c(4:6), c(1, 6, 7, 9), c(5:8), c(1, 5, 10)),
                        list(function(oracle){}), predicted, actual,
                        validation = list("permuteFold", 2, 2))
result2 <- calcCVperformance(result2, "macro F1")

performancePlot(list(result1, result2), performanceName = "Macro F1 Score",
                title = "Comparison", boxLineColouring = "None", columnVariable = "None")

```

---

plotFeatureClasses      *Plot Density, Scatterplot, Parallel Plot or Bar Chart for Features By Class*

---

**Description**

Allows the visualisation of measurements in the data set. If targets is of type [Pairs](#), then a parallel plot is automatically drawn. If it's a single categorical variable, then a bar chart is automatically drawn.

**Usage**

```

## S4 method for signature 'matrix'
plotFeatureClasses(measurements, classes, targets, ...)
## S4 method for signature 'DataFrame'
plotFeatureClasses(measurements, classes, targets, groupBy = NULL,
  groupingName = NULL, whichNumericFeaturePlots = c("both", "density", "stripchart"),
  measurementLimits = NULL, lineWidth = 1, dotBinWidth = 1,
  xAxisLabel = NULL, yAxisLabels = c("Density", "Classes"),
  showXtickLabels = TRUE, showYtickLabels = TRUE,
  xLabelPositions = "auto", yLabelPositions = "auto",
  fontSizes = c(24, 16, 12, 12, 12),
  colours = c("#3F48CC", "#880015"), showDatasetName = TRUE, plot = TRUE)
## S4 method for signature 'MultiAssayExperiment'
plotFeatureClasses(measurements, targets, groupBy = NULL, groupingName = NULL,
  showDatasetName = TRUE, ...)

```

**Arguments**

measurements	A <a href="#">matrix</a> , <a href="#">DataFrame</a> or a <a href="#">MultiAssayExperiment</a> object containing the data. For a matrix, the rows are for features and the columns are for samples. A column with name "class" must be present in the DataFrame stored in the colData slot.
classes	Either a vector of class labels of class <a href="#">factor</a> or if the measurements are of class DataFrame a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a MultiAssayExperiment object.
targets	If measurements is a matrix or DataFrame, then a vector of numeric or character indices or the feature identifiers corresponding to the feature(s) to be plotted. If measurements is a MultiAssayExperiment, then a DataFrame of 2 columns must be specified. The first column contains the names of the tables and the second contains the names of the variables, thus each row unambiguously specifies a variable to be plotted.
groupBy	If measurements is a DataFrame, then a character vector of length 1, which contains the name of a categorical feature, may be specified. If measurements is a MultiAssayExperiment, then a character vector of length 2, which contains the name of a data table as the first element and the name of a categorical feature as the second element, may be specified. Additionally, the value "clinical" may be used to refer to the column annotation stored in the colData slot of the of the MultiAssayExperiment object. A density plot will have additional lines of different line types for each category. A strip chart plot will have a separate strip chart created for each category and the charts will be drawn in a single column on the graphics device. A parallel plot and bar chart plot will similarly be laid out.
groupingName	A label for the grouping variable to be used in plots.
...	Unused variables by the three top-level methods passed to the internal method which generates the plot(s).

<code>whichNumericFeaturePlots</code>	If the feature is a single feature and has numeric measurements, this option specifies which types of plot(s) to draw. The default value is "both", which draws a density plot and also a strip chart below the density plot. Other options are "density" for drawing only a density plot and "stripchart" for drawing only a strip chart.
<code>measurementLimits</code>	The minimum and maximum expression values to plot. Default: NULL. By default, the limits are automatically computed from the data values.
<code>lineWidth</code>	Numeric value that alters the line thickness for density plots. Default: 1.
<code>dotBinWidth</code>	Numeric value that alters the diameter of dots in the strip chart. Default: 1.
<code>xAxisLabel</code>	The axis label for the plot's horizontal axis. Default: NULL.
<code>yAxisLabels</code>	A character vector of length 1 or 2. If the feature's measurements are numeric and <code>whichNumericFeaturePlots</code> has the value "both", the first value is the y-axis label for the density plot and the second value is the y-axis label for the strip chart. Otherwise, if the feature's measurements are numeric and only one plot is drawn, then a character vector of length 1 specifies the y-axis label for that particular plot. Ignored if the feature's measurements are categorical.
<code>showXtickLabels</code>	Logical. Default: TRUE. If set to FALSE, the x-axis labels are hidden.
<code>showYtickLabels</code>	Logical. Default: TRUE. If set to FALSE, the y-axis labels are hidden.
<code>xLabelPositions</code>	Either "auto" or a vector of values. The positions of labels on the x-axis. If "auto", the placement of labels is automatically calculated.
<code>yLabelPositions</code>	Either "auto" or a vector of values. The positions of labels on the y-axis. If "auto", the placement of labels is automatically calculated.
<code>fontSizes</code>	A vector of length 5. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values. The fourth number is the size of the legends' titles. The fifth number is the font size of the legend labels.
<code>colours</code>	The colours to plot data of each class in. The length of this vector must be as long as the distinct number of classes in the data set.
<code>showDatasetName</code>	Logical. Default: TRUE. If TRUE and the data is in a <code>MultiAssayExperiment</code> object, the the name of the table in which the feature is stored in is added to the plot title.
<code>plot</code>	Logical. Default: TRUE. If TRUE, a plot is produced on the current graphics device.

### Value

Plots are created on the current graphics device and a list of plot objects is invisibly returned. The classes of the plot object are determined based on the type of data plotted and the number of plots per feature generated. If the plotted variable is discrete or if the variable is numeric and one plot



type was specified, the list element is an object of class `ggplot`. Otherwise, if the variable is numeric and both the density and stripchart plot types were made, the list element is an object of class `TableGrob`.

Settling `lineWidth` and `dotBinWidth` to the same value doesn't result in the density plot and the strip chart having elements of the same size. Some manual experimentation is required to get similarly sized plot elements.

### Author(s)

Dario Strbenac

### Examples

```
# First 25 samples and first 5 genes are mixtures of two normals. Last 25 samples are
# one normal.
genesMatrix <- sapply(1:15, function(geneColumn) c(rnorm(5, 5, 1)))
genesMatrix <- cbind(genesMatrix, sapply(1:10, function(geneColumn) c(rnorm(5, 15, 1))))
genesMatrix <- cbind(genesMatrix, sapply(1:25, function(geneColumn) c(rnorm(5, 9, 2))))
genesMatrix <- rbind(genesMatrix, sapply(1:50, function(geneColumn) rnorm(95, 9, 3)))
rownames(genesMatrix) <- paste("Gene", 1:100)
colnames(genesMatrix) <- paste("Sample", 1:50)
classes <- factor(rep(c("Poor", "Good"), each = 25), levels = c("Good", "Poor"))
plotFeatureClasses(genesMatrix, classes, targets = "Gene 4",
                   xAxisLabel = bquote(log[2]*'(expression)'), dotBinWidth = 0.5)

infectionResults <- c(rep(c("No", "Yes"), c(20, 5)), rep(c("No", "Yes"), c(5, 20)))
genders <- factor(rep(c("Male", "Female"), each = 10, length.out = 50))
clinicalData <- DataFrame(Gender = genders, Sugar = runif(50, 4, 10),
                        Infection = factor(infectionResults, levels = c("No", "Yes")),
                        row.names = colnames(genesMatrix))
plotFeatureClasses(clinicalData, classes, targets = "Infection")
plotFeatureClasses(clinicalData, classes, targets = "Infection", groupBy = "Gender")

dataContainer <- MultiAssayExperiment(list(RNA = genesMatrix),
                                         colData = cbind(clinicalData, class = classes))
targetFeatures <- DataFrame(table = "RNA", feature = "Gene 50")
plotFeatureClasses(dataContainer, targets = targetFeatures,
                   groupBy = c("clinical", "Gender"),
                   xAxisLabel = bquote(log[2]*'(expression)'))
```

### Description

Collects the function to be used for making predictions and any associated parameters.

The function specified must return either a factor vector of class predictions, or a numeric vector of scores for the second class, according to the levels of the class vector of the input data set, or a data frame which has two columns named class and score.

### Constructor

`PredictParams()` Creates a default `PredictParams` object. This assumes that the object returned by the classifier has a list element named "class".

`PredictParams(predictor, intermediate = character(0), ...)` Creates a `PredictParams` object which stores the function which will do the class prediction, if required, and parameters that the function will use. If the training function also makes predictions, this must be set to `NULL`.

`predictor` Either `NULL` or a function to make predictions with. If it is a function, then the first argument must accept the classifier made in the training step. The second argument must accept a `DataFrame` of new data.

`intermediate` Character vector. Names of any variables created in prior stages in `runTest` that need to be passed to the prediction function.

... Other arguments that predictor may use.

### Author(s)

Dario Strbenac

### Examples

```
predictParams <- PredictParams(predictor = DLDApredictInterface)
# For prediction by trained object created by DLDA training function.
PredictParams(predictor = NULL)
# For when the training function also does prediction and directly returns the
# predictions.
```

---

previousSelection      *Automated Selection of Previously Selected Features*

---

### Description

Uses the feature selection of the same cross-validation iteration of a previous classification for the current classification task.

### Usage

```
## S4 method for signature 'matrix'
previousSelection(measurements, ...)
## S4 method for signature 'DataFrame'
previousSelection(measurements, classes, datasetName,
                  classifyResult, minimumOverlapPercent = 80,
                  selectionName = "Previous Selection", .iteration, verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
previousSelection(measurements, ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Do not specify this variable. It is ignored and only used to create consistency of formal parameters with other feature selection methods.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method.
datasetName	A name for the data set used. Stored in the result.
classifyResult	An existing classification result from which to take the feature selections from.
minimumOverlapPercent	If at least this many selected features can't be identified in the current data set, then the selection stops with an error.
selectionName	A name to identify this selection method by. Stored in the result.
.iteration	Do not specify this variable. It is set by <a href="#">runTests</a> if this function is being repeatedly called by <a href="#">runTests</a> .
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

**Value**

An object of class [SelectResult](#).

**Author(s)**

Dario Strbenac

**Examples**

```
#if(require(sparsediscrim))
#{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  colnames(genesMatrix) <- paste("Sample", 1:50)
  rownames(genesMatrix) <- paste("Gene", 1:100)
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  resubstitute <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "error", better = "lower")
  result <- runTests(genesMatrix, classes, datasetName = "Example",
    classificationName = "Differential Expression",
    permutations = 2, fold = 2,
    params = list(SelectParams(), TrainParams(), PredictParams()))
  features(result)

  # Genes 50 to 74 have differential expression in new data set.
  newDataset <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  newDataset <- cbind(newDataset, rbind(sapply(1:25, function(sample) rnorm(49, 9, 2)),
```

```

                                sapply(1:25, function(sample) rnorm(25, 14, 2)),
                                sapply(1:25, function(sample) rnorm(26, 9, 2)))

rownames(newDataset) <- rownames(genesMatrix)
colnames(newDataset) <- colnames(genesMatrix)

newerResult <- runTests(newDataset, classes, datasetName = "Latest Data Set",
                        classificationName = "Differential Expression",
                        permutations = 2, fold = 2,
                        params = list(SelectParams(previousSelection,
                                                  intermediate = ".iteration",
                                                  classifyResult = result),
                                     TrainParams(), PredictParams()))

# However, only genes 76 to 100 are chosen, because the feature selections are
# carried over from the first cross-validated classification.
features(newerResult)
#}

```

---

```
previousTrained
```

```
Automated Usage of Previously Created Classifiers
```

---

## Description

Uses the trained classifier of the same cross-validation iteration of a previous classification for the current classification task.

## Usage

```
## S4 method for signature 'ClassifyResult'
previousTrained(classifyResult, .iteration, verbose = 3)
```

## Arguments

classifyResult	A <a href="#">ClassifyResult</a> object which stores the models fitted previously.
.iteration	Do not specify this variable. It is set by <a href="#">runTests</a> if this function is being repeatedly called by runTests.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

## Value

A trained classifier from a previously completed classification task.

## Author(s)

Dario Strbenac

**Examples**

```

#if(require(sparsediscrim))
#{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  colnames(genesMatrix) <- paste("Sample", 1:50)
  rownames(genesMatrix) <- paste("Gene", 1:100)
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  resubstitute <- ResubstituteParams(nFeatures = seq(10, 100, 10),
    performanceType = "error", better = "lower")
  result <- runTests(genesMatrix, classes, datasetName = "Example",
    classificationName = "Differential Expression",
    permutations = 2, fold = 2,
    params = list(SelectParams(), TrainParams(), PredictParams()))

  models(result)

  # Genes 50 to 74 have differential expression in new data set.
  newDataset <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  newDataset <- cbind(newDataset, rbind(sapply(1:25, function(sample) rnorm(49, 9, 2)),
    sapply(1:25, function(sample) rnorm(25, 14, 2)),
    sapply(1:25, function(sample) rnorm(26, 9, 2))))

  rownames(newDataset) <- rownames(genesMatrix)
  colnames(newDataset) <- colnames(genesMatrix)

  newerResult <- runTests(newDataset, classes, datasetName = "Latest Data Set",
    classificationName = "Differential Expression",
    permutations = 2, fold = 2,
    params = list(SelectParams(previousSelection,
      intermediate = ".iteration",
      classifyResult = result),
      TrainParams(previousTrained,
        intermediate = ".iteration",
        classifyResult = result),
      PredictParams()))

  models(newerResult)
#}

```

---

randomForest

*Trained randomForest Object*


---

**Description**

Enables S4 method dispatching on it.

**Author(s)**

Dario Strbenac

---

 randomForestInterface *An Interface for randomForest Package's randomForest Function*


---

**Description**

A random forest classifier builds multiple decision trees and uses the predictions of the trees to determine a single prediction for each test sample.

**Usage**

```
## S4 method for signature 'matrix'
randomForestTrainInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
randomForestTrainInterface(measurements, classes, ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
randomForestTrainInterface(measurements, targets = names(measurements), ...)
## S4 method for signature 'randomForest,matrix'
randomForestPredictInterface(forest, test, ...)
## S4 method for signature 'randomForest,DataFrame'
randomForestPredictInterface(forest, test, ...,
                             returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'randomForest,MultiAssayExperiment'
randomForestPredictInterface(forest, test, targets = names(test), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
forest	A trained random forest classifier, as created by <a href="#">randomForestTrainInterface</a> , which has the same form as the output of <a href="#">randomForest</a> .
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that integer variables from the clinical data table will be used.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method (e.g. <code>verbose</code> ) or options which are accepted by the <a href="#">randomForest</a> or <a href="#">predict.randomForest</a> functions.

returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, score for a sample belonging to the second class, as determined by the factor levels, or both labels and scores in a data.frame.
verbose	Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

If measurements is an object of class MultiAssayExperiment, the factor of sample classes must be stored in the DataFrame accessible by the colData function with column name "class".

### Value

For randomForestTrainInterface, the trained random forest. For randomForestPredictInterface, either a factor vector of predicted classes, a matrix of scores for each class, or a table of both the class labels and class scores, depending on the setting of returnType.

### Author(s)

Dario Strbenac

### Examples

```
if(require(randomForest))
{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix), sep = ' ')
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix), sep = '-')
  trainingSamples <- c(1:20, 26:45)
  testingSamples <- c(21:25, 46:50)

  trained <- randomForestTrainInterface(genesMatrix[, trainingSamples],
    classes[trainingSamples])
  predicted <- randomForestPredictInterface(trained, genesMatrix[, testingSamples])
}
```

## Description

Pair-wise overlaps can be done for two types of analyses. Firstly, each cross-validation iteration can be considered within a single classification. This explores the feature ranking stability. Secondly, the overlap may be considered between different classification results. This approach compares the feature ranking commonality between different methods. Two types of commonality are possible to analyse. One summary is the average pair-wise overlap between a level of the comparison factor and the other summary is the pair-wise overlap of each level of the comparison factor that is not the reference level against the reference level. The overlaps are converted to percentages and plotted as lineplots.

## Usage

```
## S4 method for signature 'list'
rankingPlot(results, topRanked = seq(10, 100, 10),
            comparison = c("within", "classificationName", "validation",
                          "datasetName", "selectionName"),
            referenceLevel = NULL,
            lineColourVariable = c("validation", "datasetName", "classificationName",
                                   "selectionName", "None"),
            lineColours = NULL, lineWidth = 1,
            pointTypeVariable = c("datasetName", "classificationName", "validation",
                                   "selectionName", "None"),
            pointSize = 2, legendLinesPointSize = 1,
            rowVariable = c("None", "datasetName", "classificationName", "validation",
                           "selectionName"),
            columnVariable = c("classificationName", "datasetName", "validation",
                               "selectionName", "None"),
            yMax = 100, fontSizes = c(24, 16, 12, 12, 12, 16),
            title = if(comparison[1] == "within") "Feature Ranking Stability" else
                    "Feature Ranking Commonality",
            xLabelPositions = seq(10, 100, 10),
            yLabel = if(is.null(referenceLevel)) "Average Common Features (%)" else
                    paste("Average Common Features with", referenceLevel, "(%)"),
            margin = grid::unit(c(1, 1, 1, 1), "lines"),
            showLegend = TRUE, plot = TRUE, parallelParams = bpparam())
```

## Arguments

results	A list of <a href="#">ClassifyResult</a> or <a href="#">SelectResult</a> objects.
topRanked	A sequence of thresholds of number of the best features to use for overlapping.
comparison	The aspect of the experimental design to compare. See <a href="#">Details</a> section for a detailed description.
referenceLevel	The level of the comparison factor to use as the reference to compare each non-reference level to. If NULL, then each level has the average pairwise overlap calculated to all other levels.
lineColourVariable	The slot name that different levels of are plotted as different line colours.



lineColours	A vector of colours for different levels of the line colouring parameter. If NULL, a default palette is used.
lineWidth	A single number controlling the thickness of lines drawn.
pointTypeVariable	The slot name that different levels of are plotted as different point shapes on the lines.
pointSize	A single number specifying the diameter of points drawn.
legendLinesPointsSize	A single number specifying the size of the lines and points in the legend, if a legend is drawn.
rowVariable	The slot name that different levels of are plotted as separate rows of lineplots.
columnVariable	The slot name that different levels of are plotted as separate columns of lineplots.
yMax	The maximum value of the percentage to plot.
fontSizes	A vector of length 6. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values. The fourth number is the size of the legends' titles. The fifth number is the font size of the legend labels. The sixth number is the font size of the titles of grouped plots, if any are produced. In other words, when rowVariable or columnVariable are not NULL.
title	An overall title for the plot.
xLabelPositions	Locations where to put labels on the x-axis.
yLabel	Label to be used for the y-axis of overlap percentages.
margin	The margin to have around the plot.
showLegend	If TRUE, a legend is plotted next to the plot. If FALSE, it is hidden.
plot	Logical. If TRUE, a plot is produced on the current graphics device.
parallelParams	An object of class <a href="#">MulticoreParam</a> or <a href="#">SnowParam</a> .

## Details

Possible values for characteristics are "datasetName", "classificationName", "selectionName", and "validation". If "None", then that graphical element is not used.

If comparison is "within", then the feature rankings are compared within a particular analysis. The result will inform how stable the feature rankings are between different iterations of cross-validation for a particular analysis. If comparison is "classificationName", then the feature rankings are compared across different classification algorithm types, for each level of "datasetName", "selectionName" and "validation". The result will inform how stable the feature rankings are between different classification algorithms, for every cross-validation scheme, selection algorithm and data set. If comparison is "selectionName", then the feature rankings are compared across different feature selection algorithms, for each level of "datasetName", "classificationName" and "validation". The result will inform how stable the feature rankings are between feature selection classification algorithms, for every data set, classification algorithm, and cross-validation scheme. If comparison is "validation", then the feature rankings are compared across different cross-validation schemes, for each level of "classificationName",

"selectionName" and "datasetName". The result will inform how stable the feature rankings are between different cross-validation schemes, for every selection algorithm, classification algorithm and every data set. If comparison is "datasetName", then the feature rankings are compared across different data sets, for each level of "classificationName", "selectionName" and "validation". The result will inform how stable the feature rankings are between different data sets, for every classification algorithm and every data set. This could be used to consider if different experimental studies have a highly overlapping feature ranking pattern.

Calculating all pair-wise set overlaps for a large cross-validation result can be time-consuming. This stage can be done on multiple CPUs by providing the relevant options to parallelParams.

### Value

An object of class ggplot and a plot on the current graphics device, if plot is TRUE.

### Author(s)

Dario Strbenac

### Examples

```

predicted <- data.frame(sample = sample(10, 100, replace = TRUE),
                        class = rep(c("Healthy", "Cancer"), each = 50))
actual <- factor(rep(c("Healthy", "Cancer"), each = 5))
features <- sapply(1:100, function(index) paste(sample(LETTERS, 3), collapse = ''))
rankList <- list(list(features[1:100], features[c(5:1, 6:100)]),
                list(features[c(1:9, 11, 10, 12:100)], features[c(1:50, 60:51, 61:100)]))
result1 <- ClassifyResult("Example", "Differential Expression",
                        "Example Selection", LETTERS[1:10], features,
                        100, rankList,
                        list(list(rankList[[1]][[1]][1:15], rankList[[1]][[2]][1:15]),
                            list(rankList[[2]][[1]][1:10], rankList[[2]][[2]][1:10])),
                        list(function(oracle){}),
                        list(predicted), actual, list("permuteFold", 2, 2))

predicted[, "class"] <- sample(predicted[, "class"])
rankList <- list(list(features[1:100], features[c(sample(20), 21:100)]),
                list(features[c(1:9, 11, 10, 12:100)], features[c(1:50, 60:51, 61:100)]))
result2 <- ClassifyResult("Example", "Differential Variability",
                        "Example Selection",
                        LETTERS[1:10], features, 100, rankList,
                        list(list(rankList[[1]][[1]][1:15], rankList[[1]][[2]][1:15]),
                            list(rankList[[2]][[1]][1:10], rankList[[2]][[2]][1:10])),
                        list(function(oracle){}),
                        list(predicted), actual, validation = list("permuteFold", 2, 2))

rankingPlot(list(result1, result2), pointTypeVariable = "classificationName")

oneRanking <- features[c(10, 8, 1, 2, 3, 4, 7, 9, 5, 6)]
otherRanking <- features[c(8, 2, 3, 4, 1, 10, 6, 9, 7, 5)]
oneResult <- SelectResult("Example", "One Method", 10, list(oneRanking), list(oneRanking[1:5]))
otherResult <- SelectResult("Example", "Another Method", 10, list(otherRanking), list(otherRanking[1:2]))

```

```
rankingPlot(list(oneResult, otherResult), comparison = "selectionName",
            referenceLevel = "One Method", topRanked = seq(2, 8, 2),
            lineColourVariable = "selectionName", columnVariable = "None",
            pointTypeVariable = "None", xLabelPositions = 1:10)
```

---

ResubstituteParams      *Parameters for Resubstitution Error Calculation*

---

## Description

Some feature selection functions provided in the framework use resubstitution error rate to choose the best number of features for classification. This class stores parameters related to that process.

## Constructor

`ResubstituteParams()` Creates a default `ResubstituteParams` object. The number of features tried is 10, 20, 30, 40, 50, 60, 70, 80, 90, 100. The performance measure used is the balanced error rate.

`ResubstituteParams(nFeatures, performanceType, better = c("lower", "higher"))` Creates a `ResubstituteParams` object, storing information about the number of top features to calculate the performance measure for, the performance measure to use, and if higher or lower values of the measure are better.

`nFeatures` A vector for the top number of features to test the resubstitution error rate for.

`performanceType` One of the eleven types of performance metrics which can be calculated by [calcCVperformance](#).

`better` Either "lower" or "higher". Determines whether higher or lower values of the performance measure are desirable.

`intermediate` Character vector. Names of any variables created in prior stages by [runTest](#) that need to be passed to the classifier.

... Other named parameters which will be used by the classifier.

## Author(s)

Dario Strbenac

## Examples

```
ResubstituteParams(nFeatures = seq(25, 1000, 25), performanceType = "error", better = "lower")
```

ROCplot

*Plot Receiver Operating Curve Graphs for Classification Results***Description**

Creates one ROC plot or multiple ROC plots for a list of ClassifyR objects. One plot is created if the data set has two classes and multiple plots are created if the data set has three or more classes.

**Usage**

```
## S4 method for signature 'list'
ROCplot(results, nBins = sapply(results, totalPredictions),
  comparisonVariable = c("classificationName", "datasetName", "selectionName",
    "validation", "None"), lineColours = NULL,
  lineWidth = 1, fontSizes = c(24, 16, 12, 12, 12), labelPositions = seq(0.0, 1.0, 0.2),
  plotTitle = "ROC", legendTitle = NULL, xLabel = "False Positive Rate",
  yLabel = "True Positive Rate", plot = TRUE, showAUC = TRUE)
```

**Arguments**

results	A list of <a href="#">ClassifyResult</a> objects.
nBins	The number of intervals to group the samples' scores into. By default, there are as many bins as there were predictions made, for each result object.
comparisonVariable	If the data set has two classes, then the slot name with factor levels to be used for colouring the lines. Possible values are "datasetName", "classificationName", and "validation". If "None", then all lines drawn will be black. Otherwise, it specifies the variable used for plot facetting.
lineColours	A vector of colours for different levels of the line colouring parameter. If NULL, a default colour palette is used.
lineWidth	A single number controlling the thickness of lines drawn.
fontSizes	A vector of length 5. The first number is the size of the title. The second number is the size of the axes titles and AUC text, if it is not part of the legend. The third number is the size of the axes values. The fourth number is the size of the legends' titles. The fifth number is the font size of the legend labels.
labelPositions	Locations where to put labels on the x and y axes.
plotTitle	An overall title for the plot.
legendTitle	A default name is used if the value is NULL. Otherwise a character name can be provided.
xLabel	Label to be used for the x-axis of false positive rate.
yLabel	Label to be used for the y-axis of true positive rate.
plot	Logical. If TRUE, a plot is produced on the current graphics device.
showAUC	Logical. If TRUE, the AUC value of each result is added to its legend text.

## Details

The scores stored in the results should be higher if the sample is more likely to be from the class which the score is associated with. The score for each class must be in a column which has a column name equal to the class name.

For cross-validated classification, all predictions from all iterations are considered simultaneously, to calculate one curve per classification.

The number of bins determines how many pairs of TPR and FPR points will be used to draw the plot. A higher number will result in a smoother ROC curve.

The AUC is calculated using the trapezoidal rule.

## Value

An object of class `ggplot` and a plot on the current graphics device, if `plot` is `TRUE`.

## Author(s)

Dario Strbenac

## Examples

```
predicted <- list(data.frame(sample = LETTERS[c(1, 8, 15, 3, 11, 20, 19, 18)],
  Healthy = c(0.89, 0.68, 0.53, 0.76, 0.13, 0.20, 0.60, 0.25),
  Cancer = c(0.11, 0.32, 0.47, 0.24, 0.87, 0.80, 0.40, 0.75)),
  data.frame(sample = LETTERS[c(11, 18, 15, 4, 6, 10, 11, 12)],
  Healthy = c(0.45, 0.56, 0.33, 0.56, 0.33, 0.20, 0.60, 0.40),
  Cancer = c(0.55, 0.44, 0.67, 0.44, 0.67, 0.80, 0.40, 0.60)))
actual <- factor(c(rep("Healthy", 10), rep("Cancer", 10)), levels = c("Healthy", "Cancer"))
result1 <- ClassifyResult("Example", "Differential Expression", "t-test",
  LETTERS[1:20], LETTERS[10:1], 100,
  list(1:100, c(1:9, 11:101)), list(sample(10, 10), sample(10, 10)),
  list(function(oracle){}), predicted, actual,
  list("permuteFold", 2, 2))

predicted[[1]][c(2, 6), "Healthy"] <- c(0.40, 0.60)
predicted[[1]][c(2, 6), "Cancer"] <- c(0.60, 0.40)
result2 <- ClassifyResult("Example", "Differential Variability", "Bartlett Test",
  LETTERS[1:20], LETTERS[10:1], 100, list(1:100, c(1:5, 11:105)),
  list(sample(10, 10), sample(10, 10)), list(function(oracle){}),
  predicted, actual, validation = list("permuteFold", 2, 2))
ROCplot(list(result1, result2), comparisonVariable = "classificationName",
  plotTitle = "Cancer ROC")
```

---

runTest

*Perform a Single Classification*

---

## Description

For a data set of features and samples, the classification process is run. It consists of data transformation, feature selection, classifier training and testing.

**Usage**

```

## S4 method for signature 'matrix'
runTest(measurements, classes, ...)
## S4 method for signature 'DataFrame'
runTest(measurements, classes,
        balancing = c("downsample", "upsample", "none"), featureSets = NULL, metaFeatures = NULL,
        minimumOverlapPercent = 80, datasetName, classificationName,
        training, testing, params = list(SelectParams(), TrainParams(), PredictParams()),
        verbose = 1, .iteration = NULL)
## S4 method for signature 'MultiAssayExperiment'
runTest(measurements, targets = names(measurements), ...)
## S4 method for signature 'MultiAssayExperiment'
runTestEasyHard(measurements, balancing = c("downsample", "upsample", "none"),
                easyDatasetID = "clinical", hardDatasetID = names(measurements)[1],
                featureSets = NULL, metaFeatures = NULL, minimumOverlapPercent = 80,
                datasetName = NULL, classificationName = "Easy-Hard Classifier", training, testing, ..., v

```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples. The sample identifiers must be present as column names of the matrix or the row names of the DataFrame.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement object is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
balancing	A character vector of length 1 specifying how to balance samples in the training set so that each class ends up with an equal number of samples. "downsample" samples from each class with more samples than the smallest class so that each of those classes ends up with the same number of samples as the smallest class. "upsample" causes sampling with replacement to be done for every class that has less samples than the largest class to make all classes have equal size. Lastly, "none" means that no downsampling or upsampling is done and the class imbalance is preserved.
featureSets	An object of type <a href="#">FeatureSetCollection</a> which defines sets of features or sets of edges.
metaFeatures	Either NULL or a <a href="#">DataFrame</a> which has meta-features of the numeric data of interest.
minimumOverlapPercent	If featureSets stores sets of features, the minimum overlap of feature IDs with measurements for a feature set to be retained in the analysis. If featureSets stores sets of network edges, the minimum percentage of edges with both vertex IDs found in measurements that a set has to have to be retained in the analysis.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.



```

        PredictParams(DLDApredictInterface)
    ),
    training = (1:ncol(measurements)) %% 2 == 0,
    testing = (1:ncol(measurements)) %% 2 != 0)
#}

genesMatrix <- matrix(c(rnorm(90, 9, 1),
                       9.5, 9.4, 5.2, 5.3, 5.4, 9.4, 9.6, 9.9, 9.1, 9.8),
                      ncol = 10, byrow = TRUE)

colnames(genesMatrix) <- paste("Sample", 1:10)
rownames(genesMatrix) <- paste("Gene", 1:10)
genders <- factor(c("Male", "Male", "Female", "Female", "Female",
                   "Female", "Female", "Female", "Female", "Female"))

# Scenario: Male gender can predict the hard-to-classify Sample 1 and Sample 2.
clinical <- DataFrame(age = c(31, 34, 32, 39, 33, 38, 34, 37, 35, 36),
                     gender = genders,
                     class = factor(rep(c("Poor", "Good"), each = 5)),
                     row.names = colnames(genesMatrix))
dataset <- MultiAssayExperiment(ExperimentList(RNA = genesMatrix), clinical)

selParams <- SelectParams(featureSelection = differentMeansSelection, selectionName = "Difference in Means",
                          resubstituteParams = ResubstituteParams(1:10, "balanced error", "lower"))
easyHardCV <- runTestEasyHard(dataset, datasetName = "Test Data", classificationName = "Easy-Hard", training = pa
                             easyClassifierParams = list(minCardinality = 2, minPurity = 0.9),
                             hardClassifierParams = list(selParams, TrainParams(), PredictParams())
                             )

```

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runTests

*Reproducibly Run Various Kinds of Cross-Validation*


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## Description

Enables doing classification schemes such as ordinary 10-fold, 100 permutations 5-fold, and leave one out cross-validation. Processing in parallel is possible by leveraging the package [BiocParallel](#).

Pre-validation is possible and activated by specifying a list named "prevalidated" to params, which will use the functions specified in the list of parameters on the pre-validated data table. Other named items in the list correspond to other assays to be added as a pre-validated vector of the clinical table.

## Usage

```

## S4 method for signature 'matrix'
runTests(measurements, classes, ...)
## S4 method for signature 'DataFrame'
runTests(measurements, classes,
         balancing = c("downsample", "upsample", "none"), featureSets = NULL, metaFeatures = NULL,
         minimumOverlapPercent = 80, datasetName, classificationName,

```



```

validation = c("permute", "leaveOut", "fold"),
permutePartition = c("fold", "split"),
permutations = 100, percent = 25, folds = 5, leave = 2,
seed, parallelParams = bparam(),
params = list(SelectParams(), TrainParams(), PredictParams()), verbose = 1)
## S4 method for signature 'MultiAssayExperiment'
runTests(measurements, targets = names(measurements), ...)
## S4 method for signature 'MultiAssayExperiment'
runTestsEasyHard(measurements, balancing = c("downsample", "upsample", "none"),
easyDatasetID = "clinical", hardDatasetID = names(measurements)[1],
featureSets = NULL, metaFeatures = NULL, minimumOverlapPercent = 80,
datasetName = NULL, classificationName = "Easy-Hard Classifier",
validation = c("permute", "leaveOut", "fold"),
permutePartition = c("fold", "split"),
permutations = 100, percent = 25, folds = 5, leave = 2,
seed, parallelParams = bparam(), ..., verbose = 1)

```

## Arguments

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a matrix, the rows are features, and the columns are samples. The sample identifiers must be present as column names of the matrix or the row names of the DataFrame. If pre-validation is activated by naming one of the lists in params "prevalidated", this variable must be of type <a href="#">MultiAssayExperiment</a> .
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class DataFrame a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a MultiAssayExperiment object.
balancing	A character vector of length 1 specifying how to balance samples in the training set so that each class ends up with an equal number of samples. "downsample" samples from each class with more samples than the smallest class so that each of those classes ends up with the same number of samples as the smallest class. "upsample" causes sampling with replacement to be done for every class that has less samples than the largest class to make all classes have equal size. Lastly, "none" means that no downsampling or upsampling is done and the class imbalance is preserved.
featureSets	An object of type <a href="#">FeatureSetCollection</a> which defines sets of features or sets of edges.
metaFeatures	Either NULL or a DataFrame which has meta-features of the numeric data of interest.
minimumOverlapPercent	If featureSets stores sets of features, the minimum overlap of feature IDs with measurements for a feature set to be retained in the analysis. If featureSets stores sets of network edges, the minimum percentage of edges with both vertex IDs found in measurements that a set has to have to be retained in the analysis.
targets	If measurements is a MultiAssayExperiment, the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.

...	For runTests, variables not used by the matrix nor the MultiAssayExperiment method which are passed into and used by the DataFrame method. For runTestsEasyHard, easyClassifierParams and hardClassifierParams to be passed to <a href="#">easyHardClassifierTrain</a>
datasetName	A name associated with the data set used.
classificationName	A name associated with the classification.
validation	Default: "permute". "permute" for repeated permuting. "leaveOut" for leaving all possible combinations of k samples as test samples. "fold" for folding of the data set (no resampling).
permutePartition	Default: "fold". Either "fold" or "split". Only applicable if validation is "permute". If "fold", then the samples are split into folds and in each iteration one is used as the test set. If "split", the samples are split into two groups, the sizes being based on the percent value. One group is used as the training set, the other is the test set.
permutations	Default: 100. Relevant when permuting is used. The number of times to do reordering of the samples before splitting or folding them.
percent	Default: 25. Used when permutation with the split method is chosen. The percentage of samples to be in the test set.
folds	Default: 5. Relevant when repeated permutations are done and permutePartition is set to "fold" or when validation is set to "fold". The number of folds to break the data set into. Each fold is used once as the test set.
leave	Default: 2. Relevant when leave-k-out cross-validation is used. The number of samples to leave for testing.
seed	The random number generator used for repeated resampling will use this seed, if it is provided. Allows reproducibility of repeated usage on the same input data.
parallelParams	An object of class <a href="#">MulticoreParam</a> or <a href="#">SnowParam</a> .
params	A list of objects of class of <a href="#">TransformParams</a> , <a href="#">SelectParams</a> , <a href="#">TrainParams</a> or <a href="#">PredictParams</a> . The order they are in the list determines the order in which the stages of classification are done in. It may also be a list of such lists for pre-validation. In that case, each list must be named and one of them must be named "prevalidated", which specifies the functions to use on the pre-validated data table.
easyDatasetID	The name of a data set in measurements or "clinical" to indicate the patient information in the column data to be used.
hardDatasetID	The name of a data set in measurements different to the value of easyDatasetID to be used for classifying the samples not classified by the easy classifier.
verbose	Default: 1. A number between 0 and 3 for the amount of progress messages to give. A higher number will produce more messages as more lower-level functions print messages.

### Value

If the predictor function made a single prediction per sample, then an object of class [ClassifyResult](#). If the predictor function made a set of predictions, then a list of such objects.

**Author(s)**

Dario Strbenac

**Examples**

```

#if(require(sparsediscrim))
#{
  data(asthma)

  resubstituteParams <- ResubstituteParams(nFeatures = seq(5, 25, 5),
                                           performanceType = "balanced error",
                                           better = "lower")
  runTests(measurements, classes, datasetName = "Asthma",
           classificationName = "Different Means", permutations = 5,
           params = list(SelectParams(differentMeansSelection, "t Statistic",
                                     resubstituteParams = resubstituteParams),
                        TrainParams(DLDAtrainInterface),
                        PredictParams(DLDApredictInterface)
           )
  )
#}

genesMatrix <- matrix(c(rnorm(90, 9, 1),
                       9.5, 9.4, 5.2, 5.3, 5.4, 9.4, 9.6, 9.9, 9.1, 9.8),
                     ncol = 10, byrow = TRUE)

colnames(genesMatrix) <- paste("Sample", 1:10)
rownames(genesMatrix) <- paste("Gene", 1:10)
genders <- factor(c("Male", "Male", "Female", "Female", "Female",
                  "Female", "Female", "Female", "Female", "Female"))

# Scenario: Male gender can predict the hard-to-classify Sample 1 and Sample 2.
clinical <- DataFrame(age = c(31, 34, 32, 39, 33, 38, 34, 37, 35, 36),
                    gender = genders,
                    class = factor(rep(c("Poor", "Good"), each = 5)),
                    row.names = colnames(genesMatrix))
dataset <- MultiAssayExperiment(ExperimentList(RNA = genesMatrix), clinical)
selParams <- SelectParams(featureSelection = differentMeansSelection, selectionName = "Difference in Means",
                        resubstituteParams = ResubstituteParams(1:10, "balanced error", "lower"))
easyHardCV <- runTestsEasyHard(dataset, datasetName = "Test Data", classificationName = "Easy-Hard",
                              easyClassifierParams = list(minCardinality = 2, minPurity = 0.9),
                              hardClassifierParams = list(selParams, TrainParams(), PredictParams()),
                              validation = "leaveOut", leave = 1)

```

samplesMetricMap

*Plot a Grid of Sample Error Rates or Accuracies***Description**

A grid of coloured tiles is drawn. There is one column for each sample and one row for each classification result.

**Usage**

```

## S4 method for signature 'list'
samplesMetricMap(results,
  comparison = c("classificationName", "datasetName", "selectionName",
    "validation"),
  metric = c("error", "accuracy"), featureValues = NULL, featureName = NULL,
  metricColours = list(c("#3F48CC", "#6F75D8", "#9FA3E5", "#CFD1F2", "#FFFFFF"),
    c("#880015", "#A53F4F", "#C37F8A", "#E1BFC4", "#FFFFFF")),
  classColours = c("#3F48CC", "#880015"), groupColours = c("darkgreen", "yellow2"),
  fontSizes = c(24, 16, 12, 12, 12),
  mapHeight = 4, title = "Error Comparison", showLegends = TRUE,
  xAxisLabel = "Sample Name", showXtickLabels = TRUE,
  yAxisLabel = "Analysis", showYtickLabels = TRUE,
  legendSize = grid::unit(1, "lines"), plot = TRUE)

## S4 method for signature 'matrix'
samplesMetricMap(results, classes,
  metric = c("error", "accuracy"),
  featureValues = NULL, featureName = NULL,
  metricColours = list(c("#3F48CC", "#6F75D8", "#9FA3E5", "#CFD1F2", "#FFFFFF"),
    c("#880015", "#A53F4F", "#C37F8A", "#E1BFC4", "#FFFFFF")),
  classColours = c("#3F48CC", "#880015"), groupColours = c("darkgreen", "yellow2"),
  fontSizes = c(24, 16, 12, 12, 12),
  mapHeight = 4, title = "Error Comparison", showLegends = TRUE,
  xAxisLabel = "Sample Name", showXtickLabels = TRUE,
  yAxisLabel = "Analysis", showYtickLabels = TRUE,
  legendSize = grid::unit(1, "lines"), plot = TRUE)

```

**Arguments**

results	A list of <a href="#">ClassifyResult</a> objects. Could also be a matrix, for backwards compatibility.
classes	If results is a matrix, this is a factor vector of the same length as the number of columns that results has.
comparison	The aspect of the experimental design to compare.
metric	The sample-wise metric to plot.
featureValues	If not NULL, can be a named factor or named numeric vector specifying some variable of interest to plot underneath the class bar.
featureName	A label describing the information in featureValues. It must be specified if featureValues is.
metricColours	A vector of colours for metric levels.
classColours	Either a vector of colours for class levels if both classes should have same colour, or a list of length 2, with each component being a vector of the same length. The vector has the colour gradient for each class.
groupColours	A vector of colours for group levels. Only useful if groups is not NULL.
fontSizes	A vector of length 5. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values. The

	fourth number is the size of the legends' titles. The fifth number is the font size of the legend labels.
mapHeight	Height of the map, relative to the height of the class colour bar.
title	The title to place above the plot.
showLegends	Logical. IF FALSE, the legend is not drawn.
xAxisLabel	The name plotted for the x-axis. NULL suppresses label.
showXtickLabels	Logical. IF FALSE, the x-axis labels are hidden.
showYtickLabels	Logical. IF FALSE, the y-axis labels are hidden.
yAxisLabel	The name plotted for the y-axis. NULL suppresses label.
legendSize	The size of the boxes in the legends.
plot	Logical. IF TRUE, a plot is produced on the current graphics device.

### Details

The names of `results` determine the row names that will be in the plot. The length of `metricColours` determines how many bins the metric values will be discretised to.

### Value

A plot is produced and a grob is returned that can be saved to a graphics device.

### Author(s)

Dario Strbenac

### Examples

```

predicted <- data.frame(sample = LETTERS[sample(10, 100, replace = TRUE)],
                        class = rep(c("Healthy", "Cancer"), each = 50))
actual <- factor(rep(c("Healthy", "Cancer"), each = 5), levels = c("Healthy", "Cancer"))
features <- sapply(1:100, function(index) paste(sample(LETTERS, 3), collapse = ''))
result1 <- ClassifyResult("Example", "Differential Expression", "t-test",
                          LETTERS[1:10], features, 100, list(1:100), list(sample(10, 10)),
                          list(function(oracle){}), list(predicted), actual,
                          list("permuteFold", 100, 5))
predicted[, "class"] <- sample(predicted[, "class"])
result2 <- ClassifyResult("Example", "Differential Variability", "Bartlett Test",
                          LETTERS[1:10], features, 100, list(1:100), list(sample(10, 10)),
                          list(function(oracle){}), list(predicted), actual,
                          validation = list("leave", 2))
result1 <- calcCVperformance(result1, "sample error")
result2 <- calcCVperformance(result2, "sample error")
groups <- factor(rep(c("Male", "Female"), length.out = 10))
names(groups) <- LETTERS[1:10]
cholesterol <- c(4.0, 5.5, 3.9, 4.9, 5.7, 7.1, 7.9, 8.0, 8.5, 7.2)
names(cholesterol) <- LETTERS[1:10]

```

```

wholePlot <- samplesMetricMap(list(Gene = result1, Protein = result2))
wholePlot <- samplesMetricMap(list(Gene = result1, Protein = result2),
                                featureValues = groups, featureName = "Gender")
wholePlot <- samplesMetricMap(list(Gene = result1, Protein = result2),
                                featureValues = cholesterol, featureName = "Cholesterol")

```

---

selectionPlot	<i>Plot Pair-wise Overlap or Selection Size Distribution of Selected Features</i>
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### Description

Pair-wise overlaps can be done for two types of analyses. Firstly, each cross-validation iteration can be considered within a single classification. This explores the feature selection stability. Secondly, the overlap may be considered between different classification results. This approach compares the feature selection commonality between different selection methods. Two types of commonality are possible to analyse. One summary is the average pair-wise overlap between a level of the comparison factor and the other summary is the pair-wise overlap of each level of the comparison factor that is not the reference level against the reference level. The overlaps are converted to percentages and plotted as lineplots.

Additionally, a heatmap of selection size frequencies can be made.

### Usage

```

## S4 method for signature 'list'
selectionPlot(results,
              comparison = c("within", "size", "classificationName",
                            "validation", "datasetName", "selectionName"),
              referenceLevel = NULL,
              xVariable = c("classificationName", "datasetName", "validation", "selectionName"),
              boxFillColouring = c("classificationName", "size", "datasetName",
                                   "validation", "selectionName", "None"),
              boxFillColours = NULL,
              boxFillBinBoundaries = NULL, setSizeBinBoundaries = NULL,
              boxLineColouring = c("validation", "classificationName",
                                   "datasetName", "selectionName", "None"),
              boxLineColours = NULL,
              rowVariable = c("None", "validation", "datasetName",
                             "classificationName", "selectionName"),
              columnVariable = c("datasetName", "classificationName",
                                 "validation", "selectionName", "None"),
              yMax = 100, fontSizes = c(24, 16, 12, 16),
              title = if(comparison[1] == "within") "Feature Selection Stability"
                    else if(comparison == "size") "Feature Selection Size" else
                    "Feature Selection Commonality",
              xLabel = "Analysis",

```

```

yLabel = if(is.null(referenceLevel) && comparison != "size") "Common Features (%)"
          else if(comparison == "size") "Set Size" else
          paste("Common Features with", referenceLevel, "(%)"),
margin = grid::unit(c(1, 1, 1, 1), "lines"), rotate90 = FALSE,
showLegend = TRUE, plot = TRUE, parallelParams = bpparam()

```

## Arguments

results	A list of <a href="#">ClassifyResult</a> or <a href="#">SelectResult</a> objects.
comparison	The aspect of the experimental design to compare. See Details section for a detailed description.
referenceLevel	The level of the comparison factor to use as the reference to compare each non-reference level to. If NULL, then each level has the average pairwise overlap calculated to all other levels.
xVariable	The factor to make separate boxes in the boxplot for.
boxFillColouring	A factor to colour the boxes by.
boxFillColours	A vector of colours, one for each level of boxFillColouring. If NULL, a default palette is used.
boxFillBinBoundaries	Used only if comparison is "size". A vector of integers, specifying the bin boundaries of percentages of size bins observed. e.g. 0, 10, 20, 30, 40, 50.
setSizeBinBoundaries	Used only if comparison is "size". A vector of integers, specifying the bin boundaries of set size bins. e.g. 50, 100, 150, 200, 250.
boxLineColouring	A factor to colour the box lines by.
boxLineColours	A vector of colours, one for each level of boxLineColouring. If NULL, a default palette is used.
rowVariable	The slot name that different levels of are plotted as separate rows of boxplots.
columnVariable	The slot name that different levels of are plotted as separate columns of boxplots.
yMax	The maximum value of the percentage to plot.
fontSizes	A vector of length 4. The first number is the size of the title. The second number is the size of the axes titles. The third number is the size of the axes values. The fourth number is the font size of the titles of grouped plots, if any are produced. In other words, when rowVariable or columnVariable are not NULL.
title	An overall title for the plot.
xLabel	Label to be used for the x-axis.
yLabel	Label to be used for the y-axis of overlap percentages.
margin	The margin to have around the plot.
rotate90	Logical. If TRUE, the boxplot is horizontal.
showLegend	If TRUE, a legend is plotted next to the plot. If FALSE, it is hidden.
plot	Logical. If TRUE, a plot is produced on the current graphics device.
parallelParams	An object of class <a href="#">MulticoreParam</a> or <a href="#">SnowParam</a> .

## Details

Possible values for characteristics are "datasetName", "classificationName", "size", "selectionName", and "validation". If "None", then that graphical element is not used.

If comparison is "within", then the feature selection overlaps are compared within a particular analysis. The result will inform how stable the selections are between different iterations of cross-validation for a particular analysis. If comparison is "classificationName", then the feature selections are compared across different classification algorithm types, for each level of "datasetName", "selectionName" and "validation". The result will inform how stable the feature selections are between different classification algorithms, for every cross-validation scheme, selection algorithm and data set. If comparison is "selectionName", then the feature selections are compared across different feature selection algorithms, for each level of "datasetName", "classificationName" and "validation". The result will inform how stable the feature selections are between feature selection algorithms, for every data set, classification algorithm, and cross-validation scheme. If comparison is "validation", then the feature selections are compared across different cross-validation schemes, for each level of "classificationName", "selectionName" and "datasetName". The result will inform how stable the feature selections are between different cross-validation schemes, for every selection algorithm, classification algorithm and every data set. If comparison is "datasetName", then the feature selections are compared across different data sets, for each level of "classificationName", "selectionName", and "validation". The result will inform how stable the feature selections are between different data sets, for every classification algorithm and every data set. This could be used to consider if different experimental studies have a highly overlapping feature selection pattern.

Calculating all pair-wise set overlaps can be time-consuming. This stage can be done on multiple CPUs by providing the relevant options to parallelParams. The percentage is calculated as the intersection of two sets of features divided by the union of the sets, multiplied by 100.

For the selection size mode, boxFillBins is used to create bins which include the lowest value for the first bin, and the highest value for the last bin using `cut`.

## Value

An object of class ggplot and a plot on the current graphics device, if plot is TRUE.

## Author(s)

Dario Strbenac

## Examples

```
predicted <- data.frame(sample = sample(10, 100, replace = TRUE),
                        class = rep(c("Healthy", "Cancer"), each = 50))
actual <- factor(rep(c("Healthy", "Cancer"), each = 5))
features <- sapply(1:100, function(index) paste(sample(LETTERS, 3), collapse = ''))
rankList <- list(list(features[1:100], features[c(5:1, 6:100)]),
                list(features[c(1:9, 11, 10, 12:100)], features[c(1:50, 60:51, 61:100)]))
result1 <- ClassifyResult("Example", "Differential Expression",
                          "Example Selection", LETTERS[1:10], features,
                          100, rankList,
                          list(list(rankList[[1]][[1]][1:15], rankList[[1]][[2]][1:15]),
                                list(rankList[[2]][[1]][1:10], rankList[[2]][[2]][1:10])),
```



```

        list(function(oracle){}),
        list(predicted), actual, list("permuteFold", 2, 2))

predicted[, "class"] <- sample(predicted[, "class"])
rankList <- list(list(features[1:100], features[c(sample(20), 21:100)]),
                list(features[c(1:9, 11, 10, 12:100)], features[c(1:50, 60:51, 61:100)]))
result2 <- ClassifyResult("Example", "Differential Variability",
                          "Example Selection",
                          LETTERS[1:10], features, 100, rankList,
                          list(list(rankList[[1]][[1]][1:15], rankList[[1]][[2]][1:15]),
                                list(rankList[[2]][[1]][1:10], rankList[[2]][[2]][1:10])),
                          list(function(oracle){}),
                          list(predicted), actual, validation = list("permuteFold", 2, 2))

selectionPlot(list(result1, result2), xVariable = "classificationName",
              xLabel = "Analysis", columnVariable = "None", rowVariable = "None",
              boxFillColouring = "classificationName")

selectionPlot(list(result1, result2), comparison = "size",
              xVariable = "classificationName", xLabel = "Analysis",
              columnVariable = "None", rowVariable = "None",
              boxFillColouring = "size", boxFillBinBoundaries = seq(0, 100, 10),
              setSizeBinBoundaries = seq(0, 25, 5), boxLineColouring = "None")

oneRanking <- features[c(10, 8, 1, 2, 3, 4, 7, 9, 5, 6)]
otherRanking <- features[c(8, 2, 3, 4, 1, 10, 6, 9, 7, 5)]
oneResult <- SelectResult("Example", "One Method", 10, list(oneRanking), list(oneRanking[1:5]))
otherResult <- SelectResult("Example", "Another Method", 10, list(otherRanking), list(otherRanking[1:2]))

selectionPlot(list(oneResult, otherResult), comparison = "selectionName",
              xVariable = "selectionName", xLabel = "Selection Method",
              columnVariable = "None", rowVariable = "None",
              boxFillColouring = "selectionName", boxLineColouring = "None")

```

---

 SelectParams

*Parameters for Feature Selection*


---

### Description

Collects and checks necessary parameters required for feature selection. The empty constructor is provided for convenience.

### Constructor

SelectParams() Creates a default SelectParams object. This uses either an ordinary t-test or ANOVA (depending on the number of classes) and tries the top 10 to top 100 features in increments of 10, and picks the number of features with the best resubstitution balanced error rate. Users should create an appropriate SelectParams object for the characteristics of their data, once they are familiar with this software.

```
SelectParams(featureSelection, selectionName, minPresence = 1, intermediate = character(0),
             subsetToSelections = TRUE, ...)
```

Creates a `SelectParams` object which stores the function which will do the selection and parameters that the function will use.

`featureSelection` Either a function which will do the selection or a list of such functions.

For a particular function, the first argument must be an `DataFrame` object. The function's return value must be a `SelectResult` object.

`selectionName` A name to identify this selection method by.

`minPresence` If a list of functions was provided, how many of those must a feature have been selected by to be used in classification. 1 is equivalent to a set union and a number the same length as `featureSelection` is equivalent to set intersection.

`intermediate` Character vector. Names of any variables created in prior stages by `runTest` that need to be passed to a feature selection function.

`subsetToSelections` Whether to subset the data table(s), after feature selection has been done.

... Other named parameters which will be used by the selection function. If `featureSelection` was a list of functions, this must be a list of lists, as long as `featureSelection`.

#### Author(s)

Dario Strbenac

#### Examples

```
#if(require(sparsediscrim))
#{
  SelectParams(differentMeansSelection, "t-test",
              trainParams = TrainParams(), predictParams = PredictParams(),
              resubstituteParams = ResubstituteParams())

  # For pamr shrinkage selection.
  SelectParams(NSCselectionInterface, datasetName = "Cancer",
              intermediate = "trained", subsetToSelections = FALSE)
#}
```

---

SelectResult

*Container for Storing Feature Selection Results*

---

#### Description

Contains a list of ranked names of features, from most discriminative to least discriminative, and a list of features selected for use in classification. The names will be in a data frame if the input data set is a `MultiAssayExperiment`, with the first column containing the name of the data table the feature is from and the second column containing the name of the feature. Each vector or data frame element in the list corresponds to a particular iteration of classifier training. Nested lists will be present if the permutation and folding cross-validation scheme was used. This class is not intended to be created by the user, but could be used in another software package.

**Constructor**

```
SelectResult(datasetName, selectionName, totalFeatures, rankedFeatures, chosenFeatures)
```

datasetName A name associated with the data set used.

selectionName A name associated with the classification.

totalFeatures The total number of features in the data set.

rankedFeatures Identifiers of all features or meta-features if meta-features were used by the classifier, from most to least discriminative.

chosenFeatures Identifiers of features or meta-features if meta-features were used by the classifier selected at each fold.

**Summary**

A method which summarises the results is available. result is a SelectResult object.

```
show(result) Prints a short summary of what result contains.
```

**Author(s)**

Dario Strbenac

**Examples**

```
SelectResult("Asthma", "Moderated t-test", 50, list(1:50), list(1:10))
```

---

subtractFromLocation    *Subtract Numeric Feature Measurements from a Location*

---

**Description**

For each numeric feature, calculates the location, and subtracts all measurements from that location.

**Usage**

```
## S4 method for signature 'matrix'
subtractFromLocation(measurements, training, location = c("mean", "median"),
                    absolute = TRUE, verbose = 3)
## S4 method for signature 'DataFrame'
subtractFromLocation(measurements, training, location = c("mean", "median"),
                    absolute = TRUE, verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
subtractFromLocation(measurements, training, targets = names(measurements),
                    location = c("mean", "median"), absolute = TRUE, verbose = 3)
```

### Arguments

measurements	A <a href="#">matrix</a> , <a href="#">DataFrame</a> or a <a href="#">MultiAssayExperiment</a> object containing the data. For a matrix, the rows are for features and the columns are for samples.
training	A vector specifying which samples are in the training set.
location	Character. Either "mean" or "median".
absolute	Logical. Default: TRUE. If TRUE, then absolute values of the differences are returned. Otherwise, they are signed.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
verbose	Default: 3. A progress message is shown if this value is 3.

### Details

Only the samples specified by training are used in the calculation of the location. To use all samples for calculation of the location, simply provide indices of all the samples.

### Value

The same class of variable as the input variable measurements is, with the numeric features subtracted from the calculated location.

### Author(s)

Dario Strbenac

### Examples

```
aMatrix <- matrix(1:100, ncol = 10)
subtractFromLocation(aMatrix, training = 1:5, "median")
```

---

svm

*Trained svm Object*

---

### Description

Enables S4 method dispatching on it.

### Author(s)

Dario Strbenac

**Description**

SVMtrainInterface generates a trained SVM classifier and SVMpredictInterface uses it to make predictions on a test data set.

**Usage**

```
## S4 method for signature 'matrix'
SVMtrainInterface(measurements, classes, ...)
## S4 method for signature 'DataFrame'
SVMtrainInterface(measurements, classes, ..., verbose = 3)
## S4 method for signature 'MultiAssayExperiment'
SVMtrainInterface(measurements, targets = names(measurements), ...)
## S4 method for signature 'svm,matrix'
SVMpredictInterface(model, test, ...)
## S4 method for signature 'svm,DataFrame'
SVMpredictInterface(model, test, classes = NULL, returnType = c("both", "class", "score"), verbose = 3)
## S4 method for signature 'svm,MultiAssayExperiment'
SVMpredictInterface(model, test, targets = names(test), ...)
```

**Arguments**

measurements	Either a <a href="#">matrix</a> , <a href="#">DataFrame</a> or <a href="#">MultiAssayExperiment</a> containing the training data. For a <a href="#">matrix</a> , the rows are features, and the columns are samples. If of type <a href="#">DataFrame</a> , the data set is subset to only those features of type integer.
classes	Either a vector of class labels of class <a href="#">factor</a> of the same length as the number of samples in measurements or if the measurements are of class <a href="#">DataFrame</a> a character vector of length 1 containing the column name in measurement is also permitted. Not used if measurements is a <a href="#">MultiAssayExperiment</a> object.
returnType	Default: "both". Either "class", "score" or "both". Sets the return value from the prediction to either a vector of class labels, score for a sample belonging to the second class, as determined by the factor levels, or both labels and scores in a <a href="#">data.frame</a> .
test	An object of the same class as measurements with no samples in common with measurements and the same number of features as it. Also, if a <a href="#">DataFrame</a> , the class column must be absent.
targets	If measurements is a <a href="#">MultiAssayExperiment</a> , the names of the data tables to be used. "clinical" is also a valid value and specifies that numeric variables from the clinical data table will be used.
model	A fitted model as returned by SVMtrainInterface.
...	Variables not used by the <a href="#">matrix</a> nor the <a href="#">MultiAssayExperiment</a> method which are passed into and used by the <a href="#">DataFrame</a> method (e.g. verbose) or options that are used by the <a href="#">svm</a> function.

**verbose** Default: 3. A number between 0 and 3 for the amount of progress messages to give. This function only prints progress messages if the value is 3.

### Details

If `measurements` is an object of class `MultiAssayExperiment`, the factor of sample classes must be stored in the `DataFrame` accessible by the `colData` function with column name `"class"`.

### Value

For `SVMtrainInterface`, a trained SVM classifier of type `svm`. For `SVMpredictInterface`, either a factor vector of predicted classes, a vector of scores for the second class, or a table of both the class labels and second class scores, depending on the setting of `returnType`.

### Author(s)

Dario Strbenac

### Examples

```
if(require(e1071))
{
  # Genes 76 to 100 have differential expression.
  genesMatrix <- sapply(1:25, function(sample) c(rnorm(100, 9, 2)))
  genesMatrix <- cbind(genesMatrix, sapply(1:25, function(sample)
    c(rnorm(75, 9, 2), rnorm(25, 14, 2))))
  classes <- factor(rep(c("Poor", "Good"), each = 25))
  colnames(genesMatrix) <- paste("Sample", 1:ncol(genesMatrix))
  rownames(genesMatrix) <- paste("Gene", 1:nrow(genesMatrix))
  trainingSamples <- c(1:20, 26:45)
  testingSamples <- c(21:25, 46:50)

  classifier <- SVMtrainInterface(genesMatrix[, trainingSamples],
                                classes[trainingSamples], kernel = "linear")
  SVMpredictInterface(classifier, genesMatrix[, testingSamples])
}
```

### Description

Collects and checks necessary parameters required for classifier training. The empty constructor is provided for convenience.

## Constructor

`TrainParams()` Creates a default `TrainParams` object. The classifier function is DLDA. Users should create an appropriate `TrainParams` object for the characteristics of their data, once they are familiar with this software.

`TrainParams(classifier, intermediate = character(0), getFeatures = NULL, ...)`  
Creates a `TrainParams` object which stores the function which will do the classifier building and parameters that the function will use.

`classifier` A function which will construct a classifier, and also possibly make the predictions. The first argument must be a `DataFrame` object. The second argument must be a vector of classes. If the function also makes predictions and the value of the predictor setting of `PredictParams` is therefore `NULL`, the third argument must be a `DataFrame` of test data. The function must also accept a parameter named `verbose`. The function's return value can be either a trained classifier if the function only does training or a vector or data frame of class predictions if it also does prediction with the test set samples.

`intermediate` Character vector. Names of any variables created in prior stages by `runTest` that need to be passed to `classifier`.

`getFeatures` A function may be specified that extracts the selected features from the trained model. This is relevant if using a classifier that does feature selection within training (e.g. random forest). The function must return a list of two vectors. The first vector contains the ranked features (or empty if the training algorithm doesn't produce rankings) and the second vector contains the selected features.

... Other named parameters which will be used by the classifier.

## Author(s)

Dario Strbenac

## Examples

```
#if(require(sparsediscrim))
trainParams <- TrainParams(DLDAtrainInterface)
```

## Description

Collects and checks necessary parameters required for transformation. The empty constructor is for when no data transformation is desired. One data transformation function is distributed. See [subtractFromLocation](#).

**Constructor**

`TransformParams(transform, intermediate = character(0), ...)` Creates a `TransformParams` object which stores the function which will do the transformation and parameters that the function will use.

`transform` A function which will do the transformation. The first argument must be a `DataFrame` object.

`intermediate` Character vector. Names of any variables created in prior stages by `runTest` that need to be passed to a feature selection function.

`...` Other named parameters which will be used by the transformation function.

**Author(s)**

Dario Strbenac

**Examples**

```
transformParams <- TransformParams(subtractFromLocation, location = "median")  
# Subtract all values from training set median, to obtain absolute deviations.
```



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