

# Package ‘MAIT’

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**Description** The MAIT package contains functions to perform end-to-end statistical analysis of LC/MS Metabolomic Data. Special emphasis is put on peak annotation and in modular function design of the functions.

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annotateBiotransf      *Single Biotransformation Annotator*

**Description**

Function annotateBiotransf annotates a spectrum with an already detected Biotransformation.

**Usage**

```
annotateBiotransf(biotransf, diffIndex, spectrum, sigPeaksTable,biotransformationsTable)
```

**Arguments**

- biotransf      The already detected biotransformation using the [inBetween](#) function
- diffIndex      A numeric pointer to the other peak involved in the biotransformation inside the spectrum.
- spectrum      The spectrum to be annotated.

sigPeaksTable A dataframe obtained from running the [sigPeaksTable](#) function.  
 biotransformationsTable  
 Table of biotransformations either read from the bioTable argument or the default MAIT table.

### Value

A vector containing the masses of the peaks involved in the biotransformation, their retention time, their annotation and their indices in the signPeaksTable.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[Biotransformations](#)

---

Biotransformations	<i>Biotransformations of the significant data contained in a MAIT object are identified.</i>
--------------------	--

---

### Description

This function takes a [MAIT-class](#) object having significant features already detected and looks up for biotransformations between them. **MAIT** has a default biotransformation table that will be used if no other table is specified via the bioTable input parameter.

### Usage

```
Biotransformations(MAIT.object = NULL,
  peakPrecision = 0.005,
  bioTable = NULL,
  adductTable = NULL,
  adductAnnotation = FALSE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where significant features have already been found.
peakPrecision	Maximum difference between the peak masses differences and the values shown in bioTable to be considered as a biotransformation. As default the value is 0.005 Da.
bioTable	Table containing the biotransformations to be looked for in the signData input. By default it is taken the <b>MAIT-class</b> biotransformations table.
adductTable	Table containing the adducts to be looked for in the signData input. By default it is taken the <b>MAIT-class</b> positive adducts table. If this argument is set to "negAdducts", then the default table for negative adducts is taken instead. It is possible to use a user-defined adduct table

adductAnnotation

If it is set to TRUE, both adduct and Biotransformations annotation stages are performed.

**Value**

A [MAIT-class](#) object with the updated biotransformations slot

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[spectralAnova](#) [spectralTStudent](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-Biotransformations(MAIT.object = MAIT, peakPrecision = 0.005)
MAIT@FeatureInfo@biotransformations; #Detected Biotransformations
```

---

biotransformationsTable

*biotransformationsTable*

---

**Description**

This table contains the biotransformations to be looked up for.

**Value**

A table having the fields:

- NAME: The name of the biotransformation
- MASSDIFF: The mass difference of the fragment caused by the biotransformation

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[Biotransformations](#)

classes *Class names extractor from a MAIT object*

---

**Description**

Function classes extracts the class names of a linkMAIT-class object as a vector.

**Usage**

```
classes(MAIT.object)
```

**Arguments**

MAIT.object    A [MAIT-class](#) object

**Value**

A character vector containing the class names of the [MAIT-class](#) object

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT
classes(MAIT)
```

---

classifRatioClasses *Class classification ratio extractor from a MAIT object*

---

**Description**

Function classifRatioClasses extracts the class classification ratio of a MAIT object as a matrix.

**Usage**

```
classifRatioClasses(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object where function [Validation](#) has already been launched successfully.

**Value**

A matrix containing the classification ratio for each class, classifier and iteration.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT <- Validation(Iterations = 20, trainSamples= 15, MAIT.object = MAIT)
classifRatioClasses(MAIT)
```

---

classNum

*Sample number extractor for each class from a MAIT object*

---

**Description**

Function classNum extracts the number of samples belonging to each class of a MAIT object as a vector.

**Usage**

```
classNum(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

A numeric vector containing the number of samples for each class of the MAIT object. The order of the classes correspond to that of the output of function [classes](#).

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**[MAIT-class](#)**Examples**

```
data(MAIT_sample)
MAIT
classNum(MAIT)
```

---

Database

*Human Metabolome Database*

---

**Description**

The Human Metabolome Database is saved in this dataframe.

**Value**

A table having the fields:

- ENTRY: HMDB entry
- NAME: The compound name
- FORMULA: The chemical formula of the compound
- MASS: Mass of the fragment
- Biofluid: Where the compound can be found

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**[identifyMetabolites metaboliteTable](#)

---

featureID	<i>Feature ID extractor from a MAIT object</i>
-----------	--

---

**Description**

Function featureID extracts the feature IDs of a MAIT object as a vector.

**Usage**

```
featureID(MAIT.object)
```

**Arguments**

MAIT.object    A [MAIT-class](#) object

**Value**

A numeric vector containing the feature IDs of the MAIT object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
featureID(MAIT)
```

---

featureInfo	<i>Feature Info extractor from a MAIT object</i>
-------------	--

---

**Description**

Function featureInfo extracts the slot MAIT.FeatureInfo of a MAIT object.

**Usage**

```
featureInfo(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

An object of the class MAIT.FeatureInfo. More Info at [MAIT-class](#).

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
featureInfo(MAIT)
```

---

featureSigID

*Feature statistically significant ID extractor from a MAIT object*

---

**Description**

Function featureSigID extracts the vector index of the feature IDs of a MAIT object that have been found significant through function [spectralSigFeatures](#).

**Usage**

```
featureSigID(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

A numeric vector containing the statistically significant feature IDs of the MAIT object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#) [spectralSigFeatures](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
featureSigID(MAIT)
featureID(MAIT)[featureSigID(MAIT)] #Significant spectra IDs
```

---

**FisherLSD***Performs Fisher's LSD tests on the provided data*

---

**Description**

Function FisherLSD performs Fisher's LSD tests on the data using the package agricolae.

**Usage**

```
FisherLSD(data,
           classes,
           index,
           DFerror,
           MSerror,
           numClasses
           )
```

**Arguments**

data	A numerical matrix containing the data
classes	A character vector containing the class names of the samples present in the data. This vector must have the same length as the number of samples present in the argument data.
index	Numerical value to choose a subset of the data on which the LSD tests is going to be performed.
DFerror	Degrees of freedom of the model
MSerror	Means square error of the model
numClasses	Numerical parameter corresponding to the number of classes present in the data.

**Value**

A list containing the class names, the group where each class belongs according to the LSD test and the value of their means.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

---

getScoresTable	<i>Returns a list with the peak scores, masses, retention time and other information</i>
----------------	--

---

### Description

Function `getScoresTable` takes an [MAIT-class](#) object and returns a list with the scores of the features in the samples. Additionally, it returns the spectral ID of the peak and (optionally) a table containing the peak information (mass, retention time and annotation).

### Usage

```
getScoresTable(MAIT.object = NULL,
               getSpectra = TRUE,
               getExtendedTable = FALSE)
```

### Arguments

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where significant features have already been found.
<code>getSpectra</code>	If it is set to <code>TRUE</code> , an element of the returned list will contain the spectra ID of each feature.
<code>getExtendedTable</code>	If it is set to <code>TRUE</code> , an element of the returned list will contain a table with peak information (mass, retention time, annotation, intensity per sample)

### Value

A list containing:

- `scores`: The intensity of each feature per sample
- `spectraID`: A numeric with the correspondence between peaks and spectral ID
- `extendedTable`: a data frame containing detailed peak information (mass, retention time, annotation, intensity per sample).

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[spectralTStudent](#) [spectralAnova](#)

### Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
getScoresTable(MAIT,getExtendedTable=TRUE)
```

---

identifyMetabolites    *Metabolite identifier*

---

### Description

Takes a MAIT object and performs the metabolite search for the significant features

### Usage

```
identifyMetabolites(MAIT.object=NULL,  
                    peakTolerance=0.005,  
                    database=NULL,  
                    polarity="positive",  
                    printCSVfile=TRUE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where significant features have already been found.
peakTolerance	Maximum difference between the peak masses differences and the values shown in the database to be considered as a match. As default the value is 0.005 Da.
database	User-defined input table. If it is set to NULL, the default <b>MAIT</b> database is selected to perform the metabolite identification.
polarity	Character parameter that can be set to "positive" or "negative" depending on the polarity in which the samples were taken.
printCSVfile	Set to TRUE if an output table has to be produced. The table should be found in (working directory)/Tables/SearchTable.csv.

### Value

An output table is stored in the folder (working directory)/Tables/SearchTable.csv if printCSVfile is set to TRUE. More info at [metaboliteTable](#)

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[Biotransformations spectralSigFeatures](#)

### Examples

```
data(MAIT_sample)  
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)  
MAIT<-Biotransformations(MAIT.object = MAIT, peakPrecision = 0.005)  
MAIT <- identifyMetabolites(MAIT.object = MAIT, peakTolerance = 0.005,polarity="positive")
```

---

inBetween	<i>Checks if a peak mass value is in a certain mass allowance window.</i>
-----------	---

---

**Description**

Function `inBetween` extracts the mass peaks of a certain spectrum provided a dataframe where the spectrum labels are in a column called `pcgroup`.

**Usage**

```
inBetween(testValue,biotRange)
```

**Arguments**

<code>testValue</code>	The peak mass value to be checked
<code>biotRange</code>	A matrix containing two numerical columns and each row refers to a certain neutral mass loss. The first column should contain the lower value (neutral mass value minus the peak allowance window) and the second column should have the higher value (neutral mass value plus the peak allowance window)

**Value**

The rows of the `biotRange` table where possible neutral losses have been detected.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[Biotransformations](#)

---

loadings	<i>Loadings extractor for either PCA or PLS models</i>
----------	--

---

**Description**

Function `loadings` returns the loading vectors for either the PCA, PLS models when functions [plotPCA](#) or [plotPLS](#) have been already respectively launched. It also can be used to retrieve the peak aggregation models.

**Usage**

```
loadings(object, type = "none", ...)
```

**Arguments**

object	A <a href="#">MAIT-class</a> object
type	A character whose value should be "PCA" or "PLS" depending on which loading vectors are wanted. If it is set to "none", the peak aggregation models are retrieved.
...	Other input

**Value**

A matrix with the loading vectors.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPCA](#) or [plotPLS](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
loadings(MAIT)

MAIT<-plotPCA(MAIT,plot3d=FALSE)
loadings(MAIT,type="PCA")

MAIT<-plotPLS(MAIT,plot3d=FALSE)
loadings(MAIT,type="PLS")
```

---

LSDResults

*Extractor of the Fisher's LSD tests from a MAIT object*

---

**Description**

Function LSDResults extracts the results of the LSD tests of a MAIT object as a matrix.

**Usage**

```
LSDResults(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

A matrix containing the results of the Fisher's LSD tests. For each row, equal letters mean that the groups are found to be equal in the test.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class FisherLSD spectralSigFeatures](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
LSDResults(MAIT)
```

---

MAIT

*MAIT*


---

**Description**

A [MAIT-class](#) object containing simulated LC/MS data

**Value**

MAIT.object     A [MAIT-class](#) object

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

---

MAIT-class

*Class "MAIT"*


---

**Description**

MAIT class objects are used in the MAIT package to perform the analysis and statistical calculus of LC/MS data. It has 5 main slots: FeatureInfo, RawData, Validation, PhenoData and FeatureData

**Slots**

FeatureInfo: Object of class [MAIT.FeatureInfo-class](#)

RawData: Object of class [MAIT.RawData-class](#)

Validation: Object of class [MAIT.Validation-class](#)

PhenoData: Object of class [MAIT.PhenoData-class](#)

FeatureData: Object of class [MAIT.FeatureData-class](#)

## Methods

**summary** signature(object = "MAIT"): This function show a summary of the workflow results performed so far including the classification results and the parameters used.

**model** signature(object = "MAIT"): returns the model for either the PCA, PLS models when functions `plotPCA` or `plotPLS` have been already respectively launched.

**scores** signature(object = "MAIT"): Retrieves the scores from a MAIT object

**loadings** signature(object = "MAIT"): Retrieves the loadings from a MAIT object

## Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

## See Also

[xsAnnotate](#)

---

MAIT.FeatureData-class

*Class "MAIT.FeatureData"*

---

## Description

MAIT.FeatureData objects are used in the MAIT package to save the feature data.

## Slots

**scores**: Here it is saved the dataset obtained after applying the [peakAggregation](#) function.

**featureID**: The ID numbers of all features are saved here.

**featureSigID**: The ID numbers of the significant features are saved here.

**LSDResults**: The results of performing a Fisher LSD test on each significant variable are saved in this slot.

**models**: The model for each feature used to obtain the scores are saved in this slot.

**pvalues**: In this slot are saved the pvalues of the features.

**pvaluesCorrection**: The pvalues corrected by multiple test correction are saved here.

**pcaModel**: PCA model generated using the function [plotPCA](#).

**plsModel**: PCA model generated using the function [plotPLS](#).

**masses**: Masses used as an input for the function [MAITbuilder](#).

**rt**: Retention time values used as an input for the function [MAITbuilder](#).

**extendedTable**: Dataframe containing the information regarding masses, retention time values, intensity and spectra IDs passes as an input for the function [MAITbuilder](#)

## Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class xsAnnotate](#)

---

MAIT.FeatureInfo-class

*Class "MAIT.FeatureInfo"*

---

**Description**

MAIT.FeatureInfo objects are used in the MAIT package to save the data related to the information of the features.

**Slots**

It stores information related to the features. It contains three extra slots:

**biotransformations:** Biotransformations found when function [Biotransformations](#) is launched.

**peakAgMethod:** In this slot is stored the table created by the function [identifyMetabolites](#). It can be retrieved quickly in R by using the function [metaboliteTable](#)

**metaboliteTable:** Peak Aggregation Method used when function [peakAggregation](#) is launched.

**Methods**

No methods defined with class "MAIT.FeatureInfo" in the signature.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class xsAnnotate](#)

---

MAIT.Parameters-class *Class "MAIT.Parameters"*

---

**Description**

This class contains all the parameters used in the MAIT run.

**Slots**

- sampleProcessing: List containing the parameters of the function [sampleProcessing](#)
- peakAnnotation: List containing the parameters of the function [peakAnnotation](#)
- peakAggregation: List containing the parameters of the function [peakAggregation](#)
- sigFeatures: List containing the parameters of the function [spectralSigFeatures](#)
- biotransformations: List containing the parameters of the function [Biotransformations](#)
- identifyMetabolites: List containing the parameters of the function [identifyMetabolites](#)
- classification: List containing the parameters of the function [Validation](#)
- plotPCA: List containing the parameters of the function [plotPCA](#)
- plotPLS: List containing the parameters of the function [plotPLS](#)
- plotHeatmap: List containing the parameters of the function [plotHeatmap](#)

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

---

MAIT.PhenoData-class    *Class "MAIT.PhenoData"*

---

**Description**

MAIT.PhenoData objects are used in the MAIT package to save the phenotype data.

**Objects from the Class**

Objects can be created by calls of the form `new("MAIT.PhenoData", ...)`.

**Slots**

The information related to the classes present in the data is stored in this slot. It has three different extra slots:

- classes: It contains the name of the classes in the data. It can be quickly accessed by using the function [classes](#)
- classNum: Vector showing the number of samples belonging to each class. It can be quickly accessed by using the function [classNum](#)
- resultsPath: In this slot is saved the direction where the project is saved. This means that all the output tables and files of the MAIT object are going to be stored in that directory. It can be quickly accessed by using the function [resultsPath](#)

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class xsAnnotate](#)

---

MAIT.RawData-class      *Class "MAIT.RawData"*

---

**Description**

MAIT.RawData objects are used in the MAIT package to save the data related to the information of the features.

**Slots**

This class contains information of the raw data and the parameters of the whole analysis. It has two slots:

**parameters:** All the parameters of the analysis are saved in this slot. It can be obtained as a matrix in R by typing `summary(parameters(MAIT.object))`

**data:** This slot contains either the `xcmsSet`-class or the [xsAnnotate](#) object, depending if the function [peakAnnotation](#) has already been launched

**Methods**

No methods defined with class "MAIT.RawData" in the signature.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class xsAnnotate](#)

---

MAIT.Validation-class *Class "MAIT.Validation"*

---

### Description

MAIT.Validation objects are used in the MAIT package to save the validation results obtained from the classification run.

### Slots

The information related to the run of the function [Validation](#) is saved here. It contains three slots:

**ovClassifRatioTable:** Summary table showing the overall classification ratios for each of the three classifiers. It can be quickly gathered by using the function [ovClassifRatioTable](#)

**ovClassifRatioData:** All the data corresponding to the overall classification ratios. It can be quickly gathered by using the function [ovClassifRatioData](#)

**classifRatioClasses:** All the data corresponding to the classification ratios per class. It can be quickly gathered by using the function [classifRatioClasses](#)

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[MAIT-class xsAnnotate](#)

---

MAITbuilder

*MAIT constructor function when using external peak data*

---

### Description

Function MAITbuilder creates a MAIT-class object for a given external data. This process allows the user to analyse external peak data through all the MAIT processing steps.

### Usage

```
MAITbuilder(data = NULL,  
            spectraID = NULL,  
            masses = NULL,  
            rt = NULL,  
            classes = NULL,  
            significantFeatures = FALSE,  
            spectraEstimation = FALSE,  
            rtRange = 0.2,  
            corThresh = 0.7)
```

**Arguments**

<code>data</code>	Matrix containing the peak intensity values for each sample. Each row should correspond to a peak and each column to a sample.
<code>spectraID</code>	Numeric corresponding to the peak spectral grouping IDs. Two peaks having the same <code>spectraID</code> means that they correspond to the same spectrum.
<code>masses</code>	Numeric that contains the masses of the peaks. It should be as long as the number of rows in the argument data.
<code>rt</code>	Numeric that contains the retention time of the peaks. It should be as long as the number of rows in the argument data.
<code>classes</code>	Character with the class labels for each sample. It should be as long as the number of columns in the argument data.
<code>significantFeatures</code>	If it is set to TRUE, all the features set as an input are considered to be significant. Functions <a href="#">Biotransformations</a> , <a href="#">identifyMetabolites</a> , <a href="#">Validation</a> , <a href="#">plotPCA</a> , <a href="#">plotPLS</a> , <a href="#">plotHeatmap</a> , <a href="#">plotBoxplot</a> are computed on the significant features only. If it is only wanted to perform an annotation process on the external peak data, this flag should be set to TRUE.
<code>spectraEstimation</code>	If it is set to TRUE, an estimation of the peak grouping into spectra is performed. This computation is based on a retention time window (set by the argument <code>rtRange</code> ) and a correlation threshold (defined by the parameter <code>corThresh</code> ).
<code>rtRange</code>	Retention time parameter used to build a window to perform an estimation of the peak grouping into spectra.
<code>corThresh</code>	Peak correlation value used to define a threshold to perform an estimation of the peak grouping into spectra.

**Value**

All the input values are stored in a new MAIT object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**Examples**

```
data(MAIT_sample)
peaks<-scores(MAIT)
aux<-getScoresTable(MAIT)
masses<-aux$extendedTable$mz
rt <- aux$extendedTable$rt
classFactor <- rep(classes(MAIT),classNum(MAIT))
importMAIT <- MAITbuilder (data=peaks,masses=masses,rt=rt,
significantFeatures=TRUE, spectraEstimation=TRUE, rtRange=0.2,
corThresh=0.7,classes=classFactor)

importMAIT
```

---

metaboliteTable	<i>Metabolite table generator</i>
-----------------	-----------------------------------

---

### Description

Takes a [MAIT-class](#) object and builds a table with the information related to the significant features and their possible identifications.

### Usage

```
metaboliteTable(MAIT.object,  
                printCSVfile = FALSE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where significant features have already been found.
printCSVfile	A boolean parameter. Set to TRUE if a csv file should be written with the metabolite table.

### Value

An output table is stored in the folder (working directory)/Tables/SearchTable.csv having the fields:

- First column: search ID number.
- Second column (mz): Peak mass.
- Third column(rt): Peak retention time (in minutes).
- The columns from the third to the column labeled "p.adj" contain number of class samples where the peak has been detected and the intensities of the peak among samples.
- The P.adjust column contains the corrected peak p-value using bonferroni.
- The p column shows the peak p-value with no multiple test correction.
- The Fisher column shows the Fisher test results for the peak. Each of the letters separated by the character "\_" corresponds to a class value. Classes having the same letters are indistinguishable whereas those having different letters are statistically different classes.
- The isotopes column shows if the peak has been identified as a possible isotope.
- The adduct column shows which kind of adduct or biotransformation could the peak be.
- Column Name contains the name of the possible metabolite identification for the peak.
- The column labeled spectra contains the spectral ID of the peak.
- Column Biofluid shows if the identified search is stored as a biofluid in the input database or not.
- The column ENTRY shows the database name of the entry for the metabolite.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[identifyMetabolites](#) [spectralAnova](#) [spectralTStudent](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-Biotransformations(MAIT.object = MAIT, peakPrecision = 0.005)
MAIT <- identifyMetabolites(MAIT.object = MAIT, peakTolerance = 0.005,polarity="positive")
head(metaboliteTable(MAIT))
```

---

method

*Peak Aggregation Method Used*

---

**Description**

Function method returns the name of the peak aggregation method used on a [MAIT-class](#) object.

**Usage**

```
method(object)
```

**Arguments**

object            A [MAIT-class](#) object

**Value**

A character with the peak aggregation method

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[peakAggregation](#)

**Examples**

```
data(MAIT_sample)
method(MAIT)
```

---

model	<i>Model extractor for either PCA or PLS models</i>
-------	---

---

### Description

Function `model` returns the model for either the PCA, PLS models when functions `plotPCA` or `plotPLS` have been already respectively launched.

### Usage

```
model(x, type)
```

### Arguments

<code>x</code>	A <a href="#">MAIT-class</a> object
<code>type</code>	A character whose value should be "PCA" or "PLS" depending on which loading vectors are wanted.

### Value

The PCA or PLS model

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[plotPCA](#) or [plotPLS](#)

### Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPCA(MAIT,plot3d=FALSE)
model(MAIT,type="PCA")

MAIT<-plotPLS(MAIT,plot3d=FALSE)
model(MAIT,type="PLS")
```

---

models	<i>Model extractor from a MAIT object</i>
--------	---

---

**Description**

Function `models` extracts the models of a MAIT object as a list.

**Usage**

```
models(MAIT.object)
```

**Arguments**

`MAIT.object` A [MAIT-class](#) object

**Value**

A list containing the models of the MAIT object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
models(MAIT)
```

---

negAdducts	<i>Negative adducts table</i>
------------	-------------------------------

---

**Description**

This table contains the adducts to be looked up for when the LC/MS polarisation mode was set to Negative. The layout of the table is that of the CAMERA adduct table.

**Value**

A table having the fields:

- ID: An ID number
- name: The adduct name
- nmol: Number of fragments in the adduct
- charge: Electric charge of the adduct
- massdiff: Mass difference in the fragment caused by the adduct
- oidscore: Numeric relating the related clusters of ions
- quasi: Binary value showing the validness of the annotation group
- ips: Four values are possible (0.25,0.5,0.75,1) depending on the likelihood of the rule

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[peakAnnotation](#)

---

ovClassifRatioData      *Overall classification ratio extractor for MAIT objects*

---

**Description**

Function ovClassifRatioData extracts the overall classification ratio for a [MAIT-class](#) object

**Usage**

```
ovClassifRatioData(MAIT.object)
```

**Arguments**

MAIT.object      A [MAIT-class](#) object

**Value**

A list containing the overall classification ratio of the [MAIT-class](#) object for each classifier.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class Validation](#)

## Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT <- Validation(Iterations = 20, trainSamples= 15, MAIT.object = MAIT)
ovClassifRatioData(MAIT)
```

---

ovClassifRatioTable    *Overall classification table extractor for MAIT objects*

---

## Description

Function ovClassifRatioData extracts the overall classification table for a [MAIT-class](#) object

## Usage

```
ovClassifRatioTable(MAIT.object)
```

## Arguments

MAIT.object    A [MAIT-class](#) object

## Value

A list containing the overall classification table of the MAIT object for each classifier showing the mean value and their standard error.

## Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

## See Also

[MAIT-class Validation](#)

## Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT <- Validation(Iterations = 20, trainSamples= 15, MAIT.object = MAIT)
ovClassifRatioTable(MAIT)
```

---

parameters	<i>Extractor of the parameters used in the whole run from a MAIT object</i>
------------	---

---

### Description

Function `parameters` extracts the slot `linkMAIT.Parameters`-class of a [MAIT-class](#) object. This class contains all the parameters that have been used in the previous functions. Typing a summary of this object, a matrix version of the parameters is obtained.

### Usage

```
parameters(MAIT.object)
```

### Arguments

`MAIT.object`     A [MAIT-class](#) object

### Value

An object of the class `MAIT.Parameters`.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[MAIT-class](#)

### Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT <- Validation(Iterations = 20, trainSamples= 15, MAIT.object = MAIT)
parameters(MAIT)
```

pcaLoadings

*Loadings extractor for the PCA model*

---

**Description**

Function `pcaLoadings` returns the loading vectors for the PCA model when function `plotPCA`

**Usage**

```
pcaLoadings(MAIT.object)
```

**Arguments**

`MAIT.object`     A [MAIT-class](#) object

**Value**

A matrix with the PCA loading vectors.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPCA](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPCA(MAIT,plot3d=FALSE)
pcaLoadings(MAIT)
```

---

pcaModel

*Model extractor for either PCA*

---

**Description**

Function `model` returns the model for PCA when function `plotPCA` have been already respectively launched.

**Usage**

```
pcaModel(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

The PCA model of the MAIT.object

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPCA](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPCA(MAIT,plot3d=FALSE)
pcaModel(MAIT)
```

---

PCApplot3d

*3D PCA scoreplots*

---

**Description**

This function takes three sets of coordinates and builds a 3D scoreplot using the package **rgl**

**Usage**

```
PCApplot3d (z,
            x,
            y,
            cols,
            axes=TRUE,
            new=TRUE)
```

**Arguments**

**z**                    A numerical vector containing the values for the z-axis.  
**y**                    A numerical vector containing the values for the y-axis.  
**x**                    A numerical vector containing the values for the x-axis.  
**axes**                Boolean parameter. Set to TRUE if axes should be plotted.  
**new**                 Boolean parameter. Set to TRUE if a new rgl plot should be created.  
**cols**                Character vector containing the colors for each sample.

**Value**

A 3D interactive plot is created using the package **rgl**

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPCA](#)

---

pcaScores

*Scores extractor for the PCA model*

---

**Description**

Function `pcaScores` returns the loading vectors for the PCA model when function [plotPCA](#)

**Usage**

```
pcaScores(MAIT.object)
```

**Arguments**

`MAIT.object`     A [MAIT-class](#) object

**Value**

A matrix with the PCA loading vectors.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPCA](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPCA(MAIT,plot3d=FALSE)
pcaScores(MAIT)
```

---

peakAggregation	<i>Performs a peak aggregation procedure to the rawData of a MAIT object</i>
-----------------	--

---

### Description

peakAggregation function applies a peak aggregation technique to the data of a [MAIT-class](#) object. Several aggregation techniques are available (see methods below).

### Usage

```
peakAggregation(MAIT.object=NULL,
                method="None",
                classes=NULL,
                samples=NULL,
                PCAscale=FALSE,
                PCAcenter=FALSE,
                scale=FALSE,
                signVariables=NULL,
                RemoveOnePeakSpectra=FALSE,
                printCSVfile=TRUE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where function <a href="#">peakAnnotation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
method	Chosen method to perform the dimensionality reduction using the non-free <b>pagR</b> package: - If it is set to "None", no reduction is performed and the spectral peaks are taken as variables. This is the default method. - If it is set to "Mean", the intensity mean value over each sample is taken and used as spectral intensity. - If it is set to "PCA", the first scores vector of a principal components analysis (PCA) decomposition is used as spectral intensity. - If it is set to "NMF", the first scores vector of a non-negative matrix factorization (NMF) is used as spectral intensity. - If it is set to "Single", the spectral peak having the highest intensity mean value over samples among all the spectral peaks is used as spectral intensity.
classes	Parameter to explicitly define the classes of the future spectralData object. If it is set to NULL this value is taken from the annotatedPeaks input.
samples	If the spectralData object has to include just a subset of the annotatedPeaks' samples, this input must be the vector having the wanted sample's IDs.
PCAscale	If method="PCA" and PCAscale is set to TRUE, then the data is scaled following the <a href="#">prcomp</a> function. If it is set to TRUE, scale input is ignored.
PCAcenter	If method="PCA" and PCAscale is set to TRUE, then the data is centered following the <a href="#">prcomp</a> function. If it is set to TRUE, scale input is ignored.

scale	If it is set to TRUE, the data is scaled through the spectral mean value. Set to FALSE by default.
signVariables	If this input field is a numeric vector, only the spectra/peaks having an ID number present in such vector are used as input data. If it is set to NULL, all the variables are taken into account to build the input data.
RemoveOnePeakSpectra	If it is set to TRUE, all the one-peak spectra are deleted from the dataSet and the resulting spectralData object will only contain spectra with more than one peak.
printCSVfile	If it is set to TRUE, an output matrix showing the spectral/peak intensity is build, where each column is a sample and each row is a variable (spectra or peak depending on the method used).

**Value**

An [MAIT-class](#) object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**Examples**

```
data(MAIT_sample)
peakAggregation(MAIT)
```

---

peakAnnotation	<i>Spectra constructor and peak annotator</i>
----------------	---

---

**Description**

peakAnnotation function performs spectra building and peak annotation using the CAMERA package on a [MAIT-class](#) object, after applying the [sampleProcessing](#) function. The resultant [xsAnnotate](#) object is stored in a [MAIT-class](#) object.

**Usage**

```
peakAnnotation(MAIT.object = NULL,
               corrWithSamp = 0.7,
               perfwhm = 0.6,
               sigma = 6,
               adductTable = NULL,
               printSpectraTable = TRUE,
               corrBetSamp = 0.75,
               pval = 0.05,
               calcIso = TRUE,
               calcCiS = TRUE,
               calcCaS = TRUE,
               graphMethod = "hcs",
               annotateAdducts = TRUE)
```

**Arguments**

MAIT.object	A <a href="#">MAIT-class</a> object where function <a href="#">sampleProcessing</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
corrWithSamp	Correlation threshold value within samples
perfwhm	This parameter is used to group two peaks depending on their retention time. Two peaks are considered to be coeluted if their retention time falls in a range defined as $Rt\_med \pm FWHM * perfwhm$ . Where $Rt\_med$ is the retention time median and FWHM is the Full Width at Half Maximum. Defined this way, perfwhm is the percentage of the width of the FWHM (Full Width at Half Maximum)
sigma	Defining the coelution range as defined in the perfwhm variable, the FWHM is obtained by the expression $FWHM = SD * sigma$ , where SD is calculated considering the peak as normally distributed.
adductTable	User-defined input table to annotate the peaks. If it is set to NULL, the default MAIT table for adducts in positive polarization is selected. If its value is "negAdducts", the default MAIT table for fragments in negative polarization is chosen. By default it is set to NULL.
printSpectraTable	If it is set to TRUE, a three-column table is build as a csv file, where the first column shows the peak mass, the second column its retention time and the third one shows its spectral ID number. This file is saved under the project directory, in the subfolder named Tables.
corrBetSamp	Correlation threshold value between samples
pval	See <a href="#">groupCorr</a> function in the <b>CAMERA</b> package
calcIso	See <a href="#">groupCorr</a> function in the <b>CAMERA</b> package
calcCiS	See <a href="#">groupCorr</a> function in the <b>CAMERA</b> package
calcCaS	See <a href="#">groupCorr</a> function in the <b>CAMERA</b> package
graphMethod	See <a href="#">groupCorr</a> function in the <b>CAMERA</b> package
annotateAdducts	If it is set to TRUE, the function will perform an adduct annotation stage.

**Value**

A [MAIT-class](#) object containing the [xsAnnotate-class](#) in the rawData slot.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[xsAnnotate](#), [xsAnnotate-class](#)

## Examples

```
#Provided that the data files are saved accordingly
#in subfolders under a folder named "data" (see vignette):
#MAIT<-sampleProcessing(dataDir = "data", project = "Results", snThres=2,rtStep=0.02)
#MAIT<-peakAnnotation(MAIT.object = MAIT,corrWithSamp = 0.7, corrBetSamp = 0.7,perfwhm = 0.6)
```

---

plotBoxplot	<i>Prints a png file for each of the significant peak/spectra present in the input</i>
-------------	--

---

## Description

This function takes a [MAIT-class](#) object containing information related to the significant features and plots a boxplot for each significant feature (peak or spectra).

## Usage

```
plotBoxplot(MAIT.object=NULL)
```

## Arguments

MAIT.object     A [MAIT-class](#) object where significant features have already been found.

## Value

A boxplot is stored as a png file for each of the significant features (peak or spectra). The files will be stored in the directory (working directory)/Boxplots

## Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

## See Also

[spectralAnova](#) [spectralTStudent](#)

## Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotBoxplot(MAIT)
```

---

plotHeatmap	<i>Builds ten heatmaps with different p-values and clustering distances</i>
-------------	---

---

### Description

This function takes a MAIT object containing information of the significant features in the data and plots 10 heatmaps. 5 different p-values (0.05, 0.01, 0.001, 1e-4 and 1e-5) and two clustering distances (euclidean and pearson) are used.

### Usage

```
plotHeatmap(MAIT.object=NULL)
```

### Arguments

MAIT.object     A [MAIT-class](#) object where significant features have already been found.

### Value

10 different heatmaps using 5 p-values (0.05, 0.01, 0.001, 1e-4 and 1e-5) and two clustering distances (euclidean and pearson) are created. The plots will be stored as png files in a folder called (working directory)/Heatmaps

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[spectralAnova](#) [spectralTStudent](#)

### Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotHeatmap(MAIT)
```

---

`plotPCA`*2D and 3D PCA scoreplots from a MAIT object*

---

### Description

This function takes a MAIT-class object containing information of the significant features in the data and performs 2D scoreplots (PC1 vs PC2, PC2 vs PC3 and PC1 vs PC3) saved as png files. Additionally it also performs an interactive 3D PCA scoreplot.

### Usage

```
plotPCA (MAIT.object=NULL,  
         Log=FALSE,  
         center=TRUE,  
         scale=TRUE,  
         plot3d=TRUE)
```

### Arguments

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where significant features have already been found.
<code>Log</code>	Set to TRUE if the data should be plotted using the logarithm of the intensity.
<code>center</code>	Set to TRUE if the data should be centered around its mean. See <a href="#">scale</a> .
<code>scale</code>	Set to TRUE if the data should be scaled. See <a href="#">scale</a> .
<code>plot3d</code>	Boolean set to TRUE if a 3D PCA scoreplot should be plot.

### Value

Three different PCA scoreplots are printed in three png files. One using PC1 vs PC2, another with PC1 vs PC3 and the last one with PC2 vs PC3. The files will be stored in the directory (working directory)/PCA\_Scoreplots. Moreover, an interactive 3D PCA scoreplot is also generated through function [PCAplot3d](#).

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[spectralAnova](#) [spectralTStudent](#)

### Examples

```
data(MAIT_sample)  
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)  
MAIT<-plotPCA(MAIT,plot3d=FALSE)
```

---

plotPLS *2D and 3D PLS scoreplots from a MAIT object*

---

### Description

This function takes a MAIT-class object containing information of the significant features in the data and performs 2D scoreplots (PC1 vs PC2, PC2 vs PC3 and PC1 vs PC3) saved as png files. Additionally it also performs an interactive 3D PLS scoreplot.

### Usage

```
plotPLS (MAIT.object=NULL,  
         Log=FALSE,  
         center=TRUE,  
         scale=TRUE,  
         plot3d=TRUE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where significant features have already been found.
Log	Set to TRUE if the data should be plotted using the logarithm of the intensity.
center	Set to TRUE if the data should be centered around its mean. See <a href="#">scale</a> .
scale	Set to TRUE if the data should be scaled. See <a href="#">scale</a> .
plot3d	Boolean set to TRUE if a 3D PCA scoreplot should be plot.

### Value

If the number of components in the PLS is found to be three or more, three different PLS scoreplots are printed in three png files. One using PC1 vs PC2, another with PC1 vs PC3 and the last one with PC2 vs PC3. If the number of components is less than three, all the possible plots of these three are created. The files will be stored in the directory (working directory)/PLS\_Scoreplots. Moreover, an interactive 3D PLS scoreplot is also generated through function [PCAplot3d](#).

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[spectralAnova](#) [spectralTStudent](#)

### Examples

```
data(MAIT_sample)  
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)  
MAIT<-plotPLS(MAIT,plot3d=FALSE)
```

---

PLSDA

*Applies PLSDA to the provided data*

---

### Description

Function PLSDA performs Fisher's LSD tests on the data using the package plsgenomics

### Usage

```
PLSDA(Xtrain,  
      Ytrain,  
      Xtest = NULL,  
      ncomp,  
      nruncv = 0,  
      alpha = 2/3,  
      priors = NULL)
```

### Arguments

Xtrain	A numerical matrix containing the data
Ytrain	A factor vector containing the class labels of the samples
Xtest	A numerical matrix containing the data whose class is to be predicted.
ncomp	Number of components to build the PCA model
nruncv	Number of cross-validation iterations to be performed for the choice of the number of latent components
alpha	The proportion of Observations to be included in the training set at each cross-validation iteration
priors	The class priors to be used for linear discriminant analysis. If unspecified, the class proportions in the training set are used.

### Value

A list containing the output of function pls.regression, the predicted class for the Xtest dataset and the number of components used.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

---

plsLoadings                      *Loadings extractor for the PLS model*

---

**Description**

Function plsLoadings returns the loading vectors for the PLS model when function [plotPLS](#)

**Usage**

```
plsLoadings(MAIT.object)
```

**Arguments**

MAIT.object      A [MAIT-class](#) object

**Value**

A matrix with the PLS loading vectors.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPLS](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPLS(MAIT,plot3d=FALSE)
plsLoadings(MAIT)
```

---

plsModel                      *Model extractor for either PLS*

---

**Description**

Function model returns the model for PLS when function [plotPLS](#) have been already respectively launched.

**Usage**

```
plsModel(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

The PLS model of the MAIT.object

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPLS](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPLS(MAIT,plot3d=FALSE)
plsModel(MAIT)
```

---

plsScores

*Scores extractor for the PLS model*

---

**Description**

Function plsScores returns the scores vectors for the PLS model when function [plotPLS](#)

**Usage**

```
plsScores(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

A matrix with the PLS loading vectors.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[plotPLS](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT<-plotPLS(MAIT,plot3d=FALSE)
plsScores(MAIT)
```

---

posAdducts	<i>Positive adducts table</i>
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**Description**

This table contains the adducts to be looked up for when the LC/MS polarisation mode was set to positive. The layout of the table is that of the CAMERA adduct table.

**Value**

A table having the fields:

- ID: An ID number
- name: The adduct name
- nmol: Number of fragments in the adduct
- charge: Electric charge of the adduct
- massdiff: Mass difference in the fragment caused by the adduct
- oidscore: Numeric relating the related clusters of ions
- quasi: Binary value showing the validness of the annotation group
- ips: Four values are possible (0.25,0.5,0.75,1) depending on the likelihood of the rule

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[peakAnnotation](#)

---

project	<i>Change the basis of the MAIT data</i>
---------	--

---

**Description**

Function `project` is used to project the data of a MAIT object to the subspace of the models generated by another MAIT object.

**Usage**

```
project(modelData,  
        projectData)
```

**Arguments**

<code>modelData</code>	The <a href="#">MAIT-class</a> object where the models to which the new data is to be projected are saved.
<code>projectData</code>	The <a href="#">MAIT-class</a> containing the data to be projected.

**Value**

A matrix containing the data contained in the `projectData` parameter already projected into the `modelData` model subspace.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

---

pvalues	<i>Pvalues extractor from a MAIT object</i>
---------	---

---

**Description**

Function `pvalues` extracts the pvalues contained in a [MAIT-class](#) object.

**Usage**

```
pvalues(MAIT.object)
```

**Arguments**

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object
--------------------------	-------------------------------------

**Value**

A numeric vector containing the pvalues of a [MAIT-class](#) object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class spectralSigFeatures](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
pvalues(MAIT)
```

---

pvaluesCorrection      *P-values correction extractor from a MAIT object*

---

**Description**

Function pvaluesCorrection returns a character showing wheter some multiple testing correction has been performed on the p-values.

**Usage**

```
pvaluesCorrection(MAIT.object)
```

**Arguments**

MAIT.object      A [MAIT-class](#) object

**Value**

The output is a character whose values could be "None" if no p-value correction has been performed or "Bonferroni" if Bonferroni multiple test correction was selected when function [spectralSigFeatures](#) was applied.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class spectralSigFeatures](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
pvaluesCorrection(MAIT)
```

---

rawData	<i>Raw data extractor from a MAIT object</i>
---------	--

---

**Description**

Function `rawData` extracts the raw data used to build the [MAIT-class](#) object

**Usage**

```
rawData(MAIT.object)
```

**Arguments**

MAIT.object     A [MAIT-class](#) object

**Value**

A list containing either a `xcmsSet` or a [xsAnnotate](#) object.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[xsAnnotate-class](#)

**Examples**

```
data(MAIT_sample)
MAIT
rawData(MAIT)
```

---

removeOnePeakSpectra	<i>Removes those spectra having just one peak</i>
----------------------	---

---

**Description**

Function `removeOnePeakSpectra` removes the spectra having just one peak

**Usage**

```
removeOnePeakSpectra(data,
  idGroup)
```

**Arguments**

data            A numerical matrix containing the peak data  
idGroup        A numeric vector containing the spectra id number of the peaks

**Value**

A peak data set without the one-peak spectra.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[peakAggregation](#)

---

resultsPath	<i>Retrieves the folder where the results are saved for a MAIT object</i>
-------------	---

---

**Description**

Function resultsPath returns the folder where the plots and tables are saved for a [MAIT-class](#) object

**Usage**

```
resultsPath(MAIT.object)
```

**Arguments**

MAIT.object    A [MAIT-class](#) object

**Value**

A character showing where the plots and tables have been stored.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)  
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)  
resultsPath(MAIT)
```

---

retrieveSpectrum      *Extractor of the mass peaks corresponding to a certain spectrum*

---

### Description

Function `retrieveSpectrum` extracts the mass peaks of a certain spectrum provided a dataframe where the spectrum labels are in a column called `pcgroup`.

### Usage

```
retrieveSpectrum(spectrumNumber, sigPeaksTable)
```

### Arguments

- `spectrumNumber` The spectrum ID number whose peaks we want to retrieve.
- `sigPeaksTable` A dataframe containing the peak data in rows. There should be a column called `pcgroup` containing the spectra correspondence for all the peaks and the first column should contain the peak masses.

### Value

A numeric vector containing the peak masses of the queried spectrum.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[Biotransformations](#)

---

sampleProcessing      *Peak detector of netCDF samples using xcms package*

---

### Description

`sampleProcessing` takes a set of `netCDF` files containing LC/MS sample data and performs a peak detection, retention time correction and peak grouping steps using the package `xcms`. A [MAIT-class](#) object is created and all the information is saved in it.

**Usage**

```

sampleProcessing(dataDir = NULL,
                snThres = 5,
                Sigma = 5/2.3548,
                mzSlices = 0.3,
                retcorrMethod = "loess",
                groupMethod = "density",
                bwGroup = 3,
                mzWidGroup = 0.25,
                filterMethod = "centWave",
                prefilter = c(3,3000),
                rtStep = 0.03,
                nSlaves = 0,
                minfrac = 0.5,
                minsamp = 1,
                peakwidth = c(5, 20),
                project = NULL,
                ppm = 10,
                family = c("gaussian", "symmetric"),
                span = 0.2,
                fwhm = 30)

```

**Arguments**

dataDir	Folder where the netCDF files are stored. The samples files must be classified in subdirectories according to their classes.
snThres	Signal to noise ratio. Setting a high value of this parameter will lead to a higher number of features although they will be more noisy.
Sigma	Standard deviation (width) of matched filtration model peak.
mzSlices	Minimum difference in m/z for peaks with overlapping retention times.
retcorrMethod	Method used to correct the retention times values of the variables.
groupMethod	Method used to build the group peaks of variables.
bwGroup	Bandwidth (standard deviation or half width at half maximum) of gaussian smoothing kernel to apply to the peak density chromatogram.
mzWidGroup	Width of overlapping m/z slices to use for creating peak density chromatograms and grouping peaks across samples.
filterMethod	Filtering method applied in the peak detection step.
prefilter	c(k, I)specifying the prefilter step for the first analysis step(ROI detection). Mass traces are only retained if they contain at least k peakswith intensity>= I.
rtStep	Step size to use for profile generation.
nSlaves	Number of slaves for parallel calculus.
project	Project folder name under which the results will be saved. This folder will be created in the working directory.
minfrac	minimum fraction of samples necessary in at least one of the sample groups for it to be a valid group. See group.density in package <b>xcms</b> for details.

minsamp	minimum number of samples necessary in at least one of the sample groups for it to be a valid group. See <code>group.density</code> in package <b>xcms</b> for details.
ppm	maximal tolerated m/z deviation in consecutive scans, in ppm (parts per million). See <code>findPeaks.centWave</code> in package <b>xcms</b> for details.
peakwidth	Chromatographic peak width, given as range (min,max) in seconds.
fwhm	See <code>fwhm</code> argument in <code>xcmsSet</code> function.
span	See <code>span</code> argument in <code>xcmsSet</code> function.
family	See <code>family</code> argument in <code>xcmsSet</code> function.

**Value**

A **MAIT-class** object containing the data of the netCDF files. The `xcmsSet-class` object can be retrieved using the function `rawData`.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**Examples**

```
#Provided that the data files are saved accordingly
#in subfolders under a folder named "data" (see vignette):
#MAIT<-sampleProcessing(dataDir = "data", project = "Results", snThres=2,rtStep=0.02)
```

---

scores	<i>Retrieves the scores from a MAIT object</i>
--------	--

---

**Description**

Function `scores` extracts the scores MAIT object

**Usage**

```
scores(object,
        type,...)
```

**Arguments**

object	A <b>MAIT-class</b> object
type	If it is set to "none", the peak aggregation models are returned. If it is set to "PCA", the PCA model is returned. If it is set to "PLS", the PCA model is returned.
...	Other input

**Value**

A numeric matrix containing the scores saved in the MAIT object

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
scores(MAIT)
```

```
MAIT<-plotPCA(MAIT,plot3d=FALSE)
scores(MAIT,type="PCA")
```

```
MAIT<-plotPLS(MAIT,plot3d=FALSE)
scores(MAIT,type="PLS")
```

---

SearchCand

*Peak search function into a database*

---

**Description**

Function SearchCand looks up for a peak into a database

**Usage**

```
SearchCand(candidate,
            dataBase,
            peakTolerance)
```

**Arguments**

candidate	The mass of the peak to be looked up into the database
dataBase	The table where the database to be used is saved. The function is build to use databases with the same layout as the MAIT's database. This database can be accessed by typing data(MAITtables) and Database.
peakTolerance	Maximum difference between the peak masses differences and the values shown in the database to be considered as a match.

**Value**

A matrix containing all the possible hits for that peak candidate

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class identifyMetabolites](#)

---

selectK	<i>Looks for the optimum number of nearest neighbours to be considered for the KNN</i>
---------	--

---

**Description**

Function selectK finds the optimum number of nearest neighbours for the K-Nearest Neighbours (KNN) algorithm.

**Usage**

```
selectK(data,  
        class,  
        max.k)
```

**Arguments**

data	A numerical matrix containing the data
class	Vector containing the class label of each sample.
max.k	Maximum number of nearest neighbours to be considered.

**Value**

A numeric value of the optimal number of neighbours to be considered in a KNN run.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[Validation](#)

---

selectPLScomp	<i>Looks for the optimum number of components to be considered for the PLSDA</i>
---------------	--

---

### Description

Function selectPLScomp finds the optimum number of components to be used by the Partial Least Squares and linear Discriminant Algorithm (PLSDA).

### Usage

```
selectPLScomp(data,  
              class,  
              max.comp)
```

### Arguments

data	A numerical matrix containing the data
class	Vector containing the class label of each sample.
max.comp	Maximum number of components to be considered.

### Value

A numeric value of the optimal number of components to be considered in a PLSDA run.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[Validation](#)

---

sigPeaksTable	<i>Build a table of the information related to the significant features contained in a MAIT object</i>
---------------	--

---

### Description

Function sigPeaksTable takes an [MAIT-class](#) object containing significant feature information and builds a table with the information related to these features.

**Usage**

```
sigPeaksTable(MAIT.object=NULL,
              printCSVfile=FALSE,
              extendedTable = TRUE,
              printAnnotation=TRUE)
```

**Arguments**

MAIT.object      A [MAIT-class](#) object where significant features have already been found.

printCSVfile    Set to TRUE if an output table has to be produced. The table should be found in (working directory)/(project directory)Tables/significativeFeatures.csv.

extendedTable   Set to TRUE the table created by the peak external data is used.

printAnnotation      Set to TRUE The peak annotation is provided in the output table

**Value**

A table containing:

- First column (mz): Peak mass
- Second column(mzmin): Minimum peak mass of the peak group.
- Third column(mzmax): Maximum peak mass of the peak group.
- Fourth column(rt): Peak retention time (in minutes).
- Fifth column(rtmin): Minimum peak retention time of the peak group.
- Sixth column(rtmax): Maximum peak retention time of the peak group.
- Seventh column(npeaks): Number of samples where the peak has been detected.
- The columns from the ninth to the column labeled "isotopes" contain number of class samples where the peak has been detected and the intensities of the peak among samples.
- The isotopes column shows if the peak has been identified as a possible isotope.
- The adduct column shows which kind of adduct could the peak be.
- The column labeled pcgroup contains the spectral ID of the peak.
- The P.adjust column contains the corrected peak p-value using post-hoc methods.
- The p column shows the peak p-value with no multiple test correction.
- The Fisher column shows the Fisher test results for the peak. Each of the letters separated by the character "\_" corresponds to a class value. Classes having the same letters are indistinguishable whereas those having different letters are statistically different classes.
- The last columns contain the mean and median values for each feature

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[spectralTStudent](#) [spectralAnova](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
head(sigPeaksTable(MAIT))
```

---

spectralAnova

*Extract significant features from a MAIT object*


---

**Description**

Function `spectralAnova` takes an [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold. The parameters of the significant features can be printed to an output table (TRUE by default).

**Usage**

```
spectralAnova(pvalue = 0.05,
             p.adj="none",
             MAIT.object = NULL,
             printCSVfile = TRUE)
```

**Arguments**

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
<code>pvalue</code>	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
<code>p.adj</code>	Post-hoc method to be used to correct the p-values.
<code>printCSVfile</code>	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.

**Value**

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#) [peakAggregation](#) [sigPeaksTable](#)

---

spectralFUN	<i>Extract significant features from a MAIT object using a user-defined test</i>
-------------	--

---

### Description

Function `spectralFUN` takes an [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold following a user-defined statistical test. The parameters of the significant features can be printed to an output table (TRUE by default).

### Usage

```
spectralFUN(pvalue=0.05,
           p.adj="none",
           MAIT.object=NULL,
           printCSVfile=TRUE,
           test.fun=NULL,
           namefun=NULL)
```

### Arguments

<code>pvalue</code>	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
<code>p.adj</code>	Post-hoc method to be used to correct the p-values.
<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
<code>printCSVfile</code>	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.
<code>test.fun</code>	Function containing the statistical test to be applied on each feature. The function should be designed to correct just one feature as the function will apply this correction to all the features in the <code>MAIT.object</code> .
<code>namefun</code>	Character with the name of the test. This name will appear in the <code>MAITparameters</code> table and in the summary of the MAIT object.

### Value

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[MAIT-class](#) [peakAggregation](#) [sigPeaksTable](#)

---

spectralKruskal	<i>Extract significant features from a MAIT object</i>
-----------------	--

---

### Description

Function `spectralKruskal` takes an [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold following a Kruskal-Wallis test. The parameters of the significant features can be printed to an output table (TRUE by default).

### Usage

```
spectralKruskal(pvalue = 0.05,  
               p.adj="none",  
               MAIT.object = NULL,  
               printCSVfile = TRUE)
```

### Arguments

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
<code>pvalue</code>	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
<code>p.adj</code>	Post-hoc method to be used to correct the p-values.
<code>printCSVfile</code>	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.

### Value

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[MAIT-class](#) [peakAggregation](#) [sigPeaksTable](#)

---

spectralSigFeatures     *Extract significant features from a MAIT object*

---

### Description

Function `spectralSigFeatures` takes a [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold. The parameters of the significant features can be printed to an output table (TRUE by default). Depending on the number of classes in the data, the function chooses between using ANOVA tests through function `spectralAnova`, or T-Student tests by using function `spectralTStudent`.

### Usage

```
spectralSigFeatures(MAIT.object = NULL,
                   pvalue = 0.05,
                   p.adj = "none",
                   printCSVfile = FALSE,
                   scale = FALSE,
                   parametric = TRUE,
                   var.equal = FALSE,
                   test.fun = NULL,
                   jitter = FALSE,
                   jitter.factor = 1,
                   jitter.amount = 0,
                   namefun = NULL)
```

### Arguments

<code>MAIT.object</code>	A <a href="#">MAIT-class</a> object where function <code>peakAggregation</code> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
<code>pvalue</code>	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
<code>p.adj</code>	Character with the name of the posthoc method to be applied to correct the pvalues. The supported methods are that of the <code>p.adjust</code> function
<code>printCSVfile</code>	Set to TRUE if an output table has to be produced. See function <code>sigPeaksTable</code> for more information.
<code>scale</code>	Set to FALSE by default. When set to TRUE, a unit variance scaling of the data when no peak aggregation is performed. If a peak aggregation method is applied, this parameter is ignored.
<code>parametric</code>	If it is set to TRUE, the statistical tests to be applied will be parametrical tests (e.g. ANOVA, TStudent or Welch's tests). Non-parametrical tests (e.g. Kruskal-Wallis, Mann-Whitney tests) are applied otherwise.
<code>var.equal</code>	Set to FALSE by default. When set to TRUE, a Student's T-Test is applied when having 2 classes in the data. If it is set to FALSE, a Welch's test is applied instead.

test.fun	Function of the user-defined posthoc method to be applied.
jitter	If it is set to TRUE, a jitter noise is added to the data. This is useful when applying Mann-Whitney tests with ties.
jitter.factor	See argument factor of the function <a href="#">jitter</a> .
jitter.amount	See argument amount of the function <a href="#">jitter</a> .
namefun	Name of the user-defined posthoc test in the argument test.fun.

**Value**

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class spectralTStudent](#) [spectralAnova](#) [sigPeaksTable](#)

**Examples**

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
```

---

spectralTStudent	<i>Extract significant features from a MAIT object for two classes</i>
------------------	--

---

**Description**

Function `spectralTStudent` takes a [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold when there only are two classes in the raw data. The parameters of the significant features can be printed to an output table (TRUE by default).

**Usage**

```
spectralTStudent(MAIT.object = NULL,
  pvalue = 0.05,
  p.adj = "none",
  printCSVfile = TRUE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
pvalue	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
p.adj	Character with the name of the posthoc method to be applied to correct the pvalues. The supported methods are that of the <a href="#">p.adjust</a> function
printCSVfile	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.

### Value

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[spectralSigFeatures](#) [sigPeaksTable](#)

---

spectralWelch

*Extract significant features from a MAIT object*

---

### Description

Function `spectralWelch` takes an [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold following a Welch test. The parameters of the significant features can be printed to an output table (TRUE by default).

### Usage

```
spectralWelch(MAIT.object = NULL,  
             pvalue = 0.05,  
             p.adj="none",  
             printCSVfile = TRUE)
```

### Arguments

MAIT.object	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
pvalue	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
p.adj	Post-hoc method to be used to correct the p-values.
printCSVfile	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.

### Value

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[MAIT-class](#) [peakAggregation](#) [sigPeaksTable](#)

---

spectralWilcox

*Extract significant features from a MAIT object*

---

### Description

Function `spectralWilcox` takes an [MAIT-class](#) object and obtains which of the variables are significant given a p-value threshold following a Mann-Witney-Wilcoxon test. The parameters of the significant features can be printed to an output table (TRUE by default).

### Usage

```
spectralWilcox(MAIT.object = NULL,  
              pvalue = 0.05,  
              p.adj="none",  
              printCSVfile = TRUE,  
              jitter = FALSE,  
              jitter.factor = 1,  
              jitter.amount = 0)
```

**Arguments**

MAIT.object	A <a href="#">MAIT-class</a> object where function <a href="#">peakAggregation</a> has already been applied. The output of the function is going to be an update of the same <a href="#">MAIT-class</a> object.
pvalue	P-value threshold. Variables having a p-value lower than this value is considered as a significant variable.
p.adj	Post-hoc method to be used to correct the p-values.
printCSVfile	Set to TRUE if an output table has to be produced. See function <a href="#">sigPeaksTable</a> for more information.
jitter	If it is set to TRUE, a jitter noise is added to the data. This is useful when applying Mann-Whitney tests with ties.
jitter.factor	See argument factor of the function <a href="#">jitter</a> .
jitter.amount	See argument amount of the function <a href="#">jitter</a> .

**Value**

A [MAIT-class](#) object containing the significant features of the scores slot of [MAIT-class](#) object used as an input.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[MAIT-class](#) [peakAggregation](#) [sigPeaksTable](#)

---

successRatio	<i>Extracts the success ratio of a truth table</i>
--------------	--

---

**Description**

Function `successRatio` extracts the success ratio (weighted ratio of samples correctly classified vs total samples) for each class and overall. The value is weighted to take into account the possible different sample number between classes.

**Usage**

```
successRatio(classes,
             tt,
             ClassWeights)
```

**Arguments**

tt	The truth table from which the success ratio should have to be extracted.
classes	Vector containing the class label of each sample.
ClassWeights	Vector containing the weights of each class

**Value**

A numeric value showing the ratio of the samples that are correctly associated according to their real classes.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

**See Also**

[Validation](#)

---

Validation	<i>Cross validated classification over the output of the function signPeaksAnova or function TStudent2Classes</i>
------------	---

---

**Description**

Function Validation performs a cross-validated classification using three different classifiers: KNN, PLSDA and SVM. The output comes in a table with the classification ratio and its standard error. The classification ratio is weighted to take into account the different sample number of each class.

**Usage**

```
Validation(Iterations=NULL,  
          MAIT.object=NULL,  
          trainSamples=NULL,  
          PCAscale=FALSE,  
          PCAcenter=TRUE,  
          RemoveOnePeakSpectra=FALSE,  
          tuneSVM=FALSE,  
          scale=TRUE)
```

**Arguments**

Iterations	Number of iterations to be performed in the classifications. For each iteration a new training group is randomly chosen.
MAIT.object	A <a href="#">MAIT-class</a> object where significant features have already been found.
trainSamples	Number of samples per class to construct the train dataset.

PCAscale	If method="PCA" and PCAscale is set to TRUE, then the data is scaled following the prcomp function. If it is set to TRUE, scale input is ignored.
PCAcenrer	If method="PCA" and PCAscale is set to TRUE, then the data is centered following the prcomp function. If it is set to TRUE, scale input is ignored.
RemoveOnePeakSpectra	If it is set to TRUE, all the one-peak spectra are deleted from the dataSet and the resulting spectralData object will only contain spectra with more than one peak.
tuneSVM	If it is set to TRUE, a tune of parameters is performed before the SVM calculus.
scale	If it is set to TRUE, the data is scaled through the spectral mean value. Set to TRUE by default.

### Value

The numerical results of the classification per class and per classifier are saved in a [MAIT-class](#) object. Additionally, a table is also included in the output both in the list (field table) and printed as a csv file in the folder (working directory)/Validation. A boxplot is also printed as a png in the same folder showing the differences between classifiers. The confusion matrices of each iteration and classifier are also stored as csv files.

### Author(s)

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

### See Also

[peakAggregation](#) [spectralAnova](#) [spectralTStudent](#) [spectralSigFeatures](#)

### Examples

```
data(MAIT_sample)
MAIT<-spectralSigFeatures(MAIT,p.adj="fdr",parametric=TRUE)
MAIT <- Validation(Iterations = 20, trainSamples= 15, MAIT.object = MAIT)
```

---

writeExcelTable      *Writes a csv table*

---

### Description

Function writeExcelTable writes a csv table with the input data.

### Usage

```
writeExcelTable(file,
                file.name)
```

**Arguments**

file                    The data to be saved in the csv file  
file.name              The name of the csv file.

**Value**

A csv file containing the data provided as input in the file parameter.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

---

writeParameterTable    *Writes a csv table containing the parameters launched in the MAIT analysis*

---

**Description**

Function writeParameterTable writes a csv table where all the provided input parameters in the whole MAIT analysis are saved.

**Usage**

```
writeParameterTable(listParameters,  
                    folder)
```

**Arguments**

listParameters    The list of parameters to be printed. This input should be an object of the class MAIT.Parameters  
folder             The folder where the csv file is going to be saved

**Value**

A csv file containing the input parameters of the whole run.

**Author(s)**

Francesc Fernandez, <francesc.fernandez.albert@upc.edu>

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