Package 'MALDIrppa'

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Type Package Title MALDI Mass Spectrometry Data Robust Pre-Processing and Analysis **Version** 1.1.0-2 Date 2024-01-23 Maintainer Javier Palarea-Albaladejo <javier.palarea@udg.edu> **Depends** R (>= 3.2.5), MALDIquant, signal, robustbase, lattice, waveslim ByteCompile yes Description Provides methods for quality control and robust preprocessing and analysis of MALDI mass spectrometry data (Palarea-Albaladejo et al. (2018) <doi:10.1093/bioinformatics/btx628>). License GPL (>= 2) **Repository** CRAN Suggests knitr, rmarkdown, testthat VignetteBuilder knitr URL https://github.com/Japal/MALDIrppa BugReports https://github.com/Japal/MALDIrppa/issues NeedsCompilation no Author Javier Palarea-Albaladejo [cre, aut] (<https://orcid.org/0000-0003-0162-669X>), Paolo Ribeca [ctb] (<https://orcid.org/0000-0001-5599-3933>) Date/Publication 2024-01-25 17:30:06 UTC

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MALDIrppa-package MALDI mass spectra robust pre-processing and analysis

Description

This package provides procedures for quality control and robust pre-processing and analysis of MALDI mass spectrometry data based on objects and methods from the MALDIquant package. Moreover, it includes some additional functionalities and data summary and management tools (see vignette).

Details

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Author(s)

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addMetadata

Description

This function adds metadata to the metaData slot of an AbstractMassObject-class class object.

Usage

```
addMetadata(x, metadata, pos)
```

Arguments

x	List of AbstractMassObject-class (MassSpectra or MassPeaks) class objects.
metadata	Vector containing the metadata to be included for each element of x (same length as x).
pos	Position of the new metadata within the metaData slot list of each element of x.

Value

List of AbstractMassObject-class class objects including the new metadata in their metaData slot.

Examples

Load example data

data(spectra) # list of MassSpectra class objects
data(type) # metadata

```
# Add metadata
```

info <- paste("Spectrum No.",1:length(spectra)) # Artificial metadata vector</pre>

```
spectra2 <- addMetadata(spectra,info,1)</pre>
```

Check info in metaData slot

```
spectra2[[1]]@metaData
```

```
alignPeaks
```

Description

This function provides a single command for selecting anchor peaks, peak alignment and binning of MassPeaks class objects (MALDIquant package). It also deals with alignment-related issues found in high-resolution mass spectrometry data.

Usage

```
alignPeaks(x, minFreq = 0.9, tolerance = 0.003, ...)
```

Arguments

х	A list of MassPeaks class objects.
minFreq	Minimum relative frequency of a peak over x to be considered as anchor peak for alignment (see referencePeaks).
tolerance	Maximal deviation in peak masses to be considered as identical (see referencePeaks, binPeaks).
	Other arguments from the original functions in MALDIquant.

Details

See warpMassPeaks and binPeaks in the MALDIquant package for details about the alignment and binning algorithms. Note that alignPeaks applies an additional binning round which helps to correct for misalignment issues found after using the default strict or relaxed bin creation rules in high-resolution mass spectrometry data.

Value

A list of MassPeaks class objects with aligned peaks along a common m/z range.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects

```
# Some pre-processing
```

```
spectra <- screenSpectra(spectra)$fspectra
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)</pre>
```

Peak alignment

countPeaks

peaks <- alignPeaks(peaks, minFreq = 0.8)</pre>

countPeaks

Count the number of peaks in MassPeaks objects

Description

This function provides the number of peaks of each element of a list of MassPeaks objects.

Usage

countPeaks(x)

Arguments

х

A list of MassPeaks objects.

Value

A vector consisting of the number of peaks for each peak profile in x.

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
# Some pre-processing
spectra <- screenSpectra(spectra)$fspectra
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
# Count peaks
```

npeaks <- countPeaks(peaks)</pre>

deletePeaks

Description

This function deletes peaks of height (intensity) below a given value in MassPeaks objects.

Usage

deletePeaks(x, min = NULL)

Arguments

Х	A list of MassPeaks objects.
min	Lower threshold used to discard a peak.

Details

This functions takes a list of MassPeaks objects and filters out peaks of height (intensity) falling below the given minimum value.

Value

A filtered list of MassPeaks objects.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects

Some pre-processing

```
spectra <- screenSpectra(spectra)$fspectra
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
# Delete peaks of intensity < 30</pre>
```

peaks <- deletePeaks(peaks, min = 30)</pre>

detectOutliers

Description

This function identifies outlying cases in a collection of processed mass peak profiles. It can be applied either on peak intensities or binary data (peak presence/absence patterns). It allows to specify a grouping factor in order to execute the procedure at the desired level of aggregation.

Usage

detectOutliers(x, by = NULL, binary = FALSE, ...)

Arguments

х	A list of MassSpectrum objects containing processed peaks.
by	If given, a grouping variable (factor or numeric) subsetting the data.
binary	Logical value. It indicates whether the procedure must be applied on either peak intensities (FALSE, default) or on binary peak presence/absence patterns (TRUE).
	Optional arguments for the robust outlier detection method.

Details

This function marks samples with mass peak profiles that largely deviates from other samples at the given aggregation level. It uses robust methods for the detection of multivariate outliers applied on metric multidimensional scaling (MDS) coordinates (Euclidean distance is used for peak intensities and binary distance for binary profiles; see dist). The number of MDS coordinates used is generally set to p = floor(n/2), where n is the number of samples in the target subset. This is an upper cap recommended for the computation of the robust MCD estimator by covMcd. However, that rule of thumb can still generate matrix singularity problems with covMcd in some cases. When this occurs detectOutliers further reduces p to use the maximum number of MDS coordinates giving rise to a non-singular covariance matrix (min(p) = 2 in any case). The adaptive multivariate outlier detection algorithm was adapted from the mvoutlier package.

Value

If by = NULL, a logical vector of length equal to the number of elements of x indicating outlying samples by TRUE. Otherwise, a 2-column data.frame is generated which includes such a logical vector along with the grouping variable given in by.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects
data(type) # metadata

```
# Some pre-processing
```

```
sc.results <- screenSpectra(spectra,meta=type)
spectra <- sc.results$fspectra # filtered mass spectra
type <- sc.results$fmeta  # filtered metadata
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
peaks <- alignPeaks(peaks, minFreq = 0.8)
# Find outlying samples at isolate level
out <- detectOutliers(peaks, by = type$isolate)
# From peak presence/absence patterns
out.binary <- detectOutliers(peaks, by = type$isolate, binary = TRUE)</pre>
```

importSpectra

Convert mass spectra from text files into MassSpectrum objects

Description

This function allows to import collections of mass spectra stored in individual text files into a list of MassSpectrum objects.

Usage

```
importSpectra(where = getwd())
```

Arguments

where

Path to the folder where the text files are stored (default: current working directory).

Details

This functions works with dat, csv or txt file types containing two columns: the first one referring to common m/z values and the second one to intensities (using single-space separator between both and no column names). It reads all the .dat, .csv or .txt files in the given folder (so unrelated files should better not be there) and creates a list of MassSpectrum objects. For importing data from more specialised file formats we refer the reader to the package MALDIquantForeign.

Value

A list of MassSpectrum objects.

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peakPatterns

Examples

```
# Create fake mass spectrometry data
s1 <- cbind(1:20, rlnorm(20))
s2 <- cbind(1:20, rlnorm(20))
s3 <- cbind(1:20, rlnorm(20))
# Save as csv files in temporary directory
path <- tempdir()
write.table(s1, file = file.path(path, "s1.csv"),
            row.names = FALSE, col.names = FALSE, sep=" ")
write.table(s2, file = file.path(path, "s2.csv"),
            row.names = FALSE, col.names = FALSE, sep=" ")
write.table(s3, file = file.path(path, "s3.csv"),
            row.names = FALSE, col.names = FALSE, sep=" ")
# Import files and arrange into a list of MassSpectrum objects
spectra <- importSpectra(where = path)</pre>
```

peakPatterns Display peak presence/absence patterns

Description

This function displays the patterns of peak presence and absence in an intensity matrix as generated from intensityMatrix.

Usage

```
peakPatterns(x, abs.lab = NA, barplot = TRUE,
    axis.lab = c("m/z", "Index"), bar.col = "red3",
    cell.col = c("white", "dodgerblue"), grid = FALSE,
    grid.col = "black", grid.lty = "dotted", cex.axis = 0.5,
    cex.lab = 0.5, ...)
```

Arguments

x	A matrix, data.frame or a list of MassPeaks objects.
abs.lab	Unique label used to denote peak absence in x (NA, default).
barplot	Logical value indicating whether a barplot of relative peak frequency across samples is displayed (TRUE, default).
axis.lab	Vector of axis labels in the c("x", "y") format.
bar.col	Colour of the bars in the barplot.

cell.col	Vector of colours for the table cells (format c("col.absence", "col.presence")).
grid	Logical value indicating whether gridlines are added (FALSE, default).
grid.col	Colour of the gridlines ("black", default).
grid.lty	Style of the gridlines ("dotted", default. See 1ty in par).
cex.axis	Axis tick labels scaling factor relative to default.
cex.lab	Axis labels scaling factor relative to default.
	Other arguments.

Details

The peak presence/absence patterns are displayed by rows from the first (top) to the last (bottom) sample in the data set x over the range of common m/z points. Positive peaks are by default represented by coloured cells whereas zero or absent peaks are left blank. A barplot on the top margins shows the relative frequency of a peak at each m/z point across samples.

Value

No return value, graphical output.

See Also

See intensityMatrix.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects
data(type) # metadata

```
# Some pre-processing
```

```
sc.results <- screenSpectra(spectra,meta=type)
spectra <- sc.results$fspectra # filtered mass spectra
type <- sc.results$fmeta # filtered metadata
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
peaks <- alignPeaks(peaks, minFreq = 0.8)</pre>
```

```
# Display patterns across all data
```

peakPatterns(peaks)

```
# Check results within isolate 280
```

```
peakPatterns(peaks[type$Isolate=="280"])
```

plot.scSpectra

Description

This is a plot method for scSpectra objects that displays the results from applying screenSpectra to identify potential faulty, low-quality raw mass spectra.

Usage

Arguments

х	A scSpectra object as generated by screenSpectra.
type	Type of graphical display.
breaks	Number of break points for plotting a histogram when type = "hist" (default = 30).
labels	Vector of labels for the mass spectra (default = FALSE, no labels).
col	Colour for the histogram bars when type = "hist".
	Other arguments.

Details

For type = "index" (default) the upper and lower fences used to declare a mass spectrum as potentially low-quality are shown along with their A scores (see screenSpectra for details). Point labels can be added using the labels argument (either a position index when labels = TRUE or a given vector of labels; see examples below). For type = "hist" a histogram of the distribution of the A scores is produced along with the tolerance fences. Finally, type = "casewise" displays interactively the flagged spectra for visual inspection.

Value

No return value, graphical output.

See Also

See screenSpectra and summary.scSpectra.

Examples

```
# Load example data
data(spectra) # list of MassSpectra objects
data(type) # metadata
sc.results <- screenSpectra(spectra)
plot(sc.results)
plot(sc.results, labels = TRUE)
plot(sc.results, labels = type$SpectID)
plot(sc.results, type = "hist")
```

rawToPeaks

Create list of MassPeaks objects

Description

This is an auxiliary function to create a list of MassPeaks objects from raw data.

Usage

rawToPeaks(mz, I)

Arguments

mz	Vector of m/z values.
I	Matrix of peak intensity values

Details

This functions creates a list of MassPeaks objects from a vector of common m/z values and a matrix of column vectors of peak intensities for a collections of mass peak profiles. The column names are used to label the elements of the list.

Value

A list of MassPeaks objects.

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rawToSpectra

Description

This is an auxiliary function to create a list of MassSpectrum objects from raw data.

Usage

```
rawToSpectra(mz, I)
```

Arguments

mz	Vector of m/z values.
I	Matrix of intensity values.

Details

This functions creates a list of MassSpectrum objects from a vector of common m/z values and a matrix of column vectors of intensities for a collections of mass spectra. The column names are used to label the elements of the list.

Value

A list of MassSpectrum objects.

See Also

See importSpectra.

redResolution Reduce resolution of MassSpectrum objects

Description

This function allows to obtain a lighter version of a list of MassSpectrum objects by decreasing their m/z resolution.

Usage

redResolution(x, by = 1)

Arguments

Х	A list of MassSpectrum objects.
by	Number of times reduction (by = 1, default).

Details

This function reduces the resolution of mass spectra by eliminating a regular sequence of m/z sampling points in steps given by the argument by. For example, specifying by = 2 means to reduce the length and memory usage of the signal by a half approximately.

Value

A list of MassSpectrum objects.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects

Reduce resolution by a half

```
spectra.LowRes <- redResolution(spectra, by = 2)</pre>
```

screenSpectra Identification of potentially low-quality raw mass spectra

Description

This function implements a quality control check to help in the identification of possibly faulty, low-quality raw mass spectra. It computes an atypicality score and labels suspicious profiles for further inspection and filtering.

Usage

Arguments

x	A list of MassSpectrum objects.
meta	(optional) Matrix or vector containing metadata associated to x. Typically a data matrix including spectrum ID, biotype, replicate number, etc. for each element of x.
threshold	Multiplicative factor used in computing the upper and lower fences to determine passes and failures. It is related to the actual method used to compute the fences (see method). Typically, threshold = 1.5 (default value) for the boxplot rules, and threshold = 3 for the others.

estimator	Robust scale estimator used:
	Q: robust location-free scale estimate (default, see Qn function in robustbase package). More efficient than MAD and adequate for non-symmetric distributions.
	MAD: median absolute deviance scale estimate. Very robust and preferred for fairly symmetric distributions.
method	Method used to compute upper and lower fences for the identification of atypical mass spectra.
	boxplot: standard boxplot rule based on the first and third quartiles and the interquartile range.
	adj.boxplot: extension of boxplot rule for strongly asymmetric data (default).
	ESD: extreme studentized deviation method. Based on the mean and the standard deviation of the data. Typically used with threshold = 3 (three-sigma rule).
	Hampel: robust version of the ESD method based on the median and the median absolute deviance estimate (MAD).
	RC: as Hampel's but replacing MAD by Rousseeuw & Croux (1993)'s Qn as scale estimate.
nd	Order for the derivative function of the mass spectra (default = 1).
lambda	Weight given to each component of the atypicality score (values in $[0, 1]$, default = 0.5, see details below).
	Other arguments.

Details

The procedure computes an atypicality score (A score) based on a weighted function of two components: (1) a robust scale estimator (Q or MAD) of the n-order derivative (computed using Savitzky-Golay smoothing filter) of scaled mass spectra and (2) the median intensity of the signals. Given a method to determine tolerance fences, a mass spectrum is labelled as potentially faulty, low-quality according to the magnitude of its A score. The adj.boxplot method based on the Q scale estimator and equal weights to both components (lambda = 0.5) are the default options. The greater lambda the higher the weight given to the scale estimator in the A score. The function produces summaries and a list of mass spectra and (if given) associated metadata in which the identified cases were filtered out.

Value

An object of class scSpectra with elements:

fspectra	List of mass spectra (MassSpectrum class) with potential low-quality cases fil- tered out.
fmeta	Associated filtered metadata (data.frame object).
est.table	Results table showing the mass spectra ID, A score and label (pass/failure).
	Other details (see method summary.scSpectra for scSpectra objects).

See Also

See methods summary.scSpectra and plot.scSpectra for scSpectra objects.

Examples

```
# Load example data
data(spectra) # list of MassSpectra objects
data(type) # metadata
# Results using different settings
sc.results <- screenSpectra(spectra)
sc.results <- screenSpectra(spectra, type)
sc.results <- screenSpectra(spectra, type, method = "RC")
sc.results <- screenSpectra(spectra, type, threshold = 3, estimator = "MAD", method = "Hampel")
# Numerical and graphical summary
summary(sc.results)
plot(sc.results)
# Save filtered data for further pre-processing
filtered.spectra <- sc.results$fspectra
filtered.type <- sc.results$fmeta</pre>
```

snrPeaks

Extract signal-to-noise ratio thresholds from MassPeaks *objects*

Description

This function extracts the thresholds used to determine peaks from mass spectra based on signal-tonoise ratio (SNR) (threshold equal to SNR*noise).

Usage

snrPeaks(x)

Arguments

Х

A list of MassPeaks objects.

Details

Given a collection of MassPeaks objects as obtained from detectPeaks, this function provides the thresholds used in each case to determine peaks from the original mass spectra. The thresholds are calculated as the product of a SNR value set by the user and the estimated noise of the signal (see detectPeaks).

Value

A list of vectors of SNR-based thresholds, one for each sample.

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spectra

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
# Some pre-processing
spectra <- screenSpectra(spectra)$fspectra
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
# Extract thresholds for each mass peak profile
SNRs <- snrPeaks(peaks)</pre>
```

spectra

Example mass spectra profiles

Description

List of mass spectra (MassSpectrum class).

Usage

data(spectra)

Details

Low-resolution version of a MALDI-TOF mass spectrometry data set in the range [2500, 13000] m/z provided for illustration purposes. It consists of 4 technical replicates of 5 biological replicates from 19 bacterial isolates (see type for associated metadata).

Examples

data(spectra)

str(spectra[[1]])

plot(spectra[[1]])

summary.scSpectra

Description

This is a summary method for scSpectra objects that generates a numerical summary of the settings and results from applying screenSpectra to identify potential faulty, low-quality raw mass spectra.

Usage

```
## S3 method for class 'scSpectra'
summary(object, ncases = 10, ...)
```

Arguments

object	A scSpectra object as generated from screenSpectra.
ncases	Number of cases shown in the results table.
	Other arguments.

Details

A table is generated that includes details of the numerical estimations along with mass spectra ID, A score and the label for each mass spectra, either potentially low-quality (failure) or good-quality (success).

Value

No return value, text printed on console.

See Also

See screenSpectra and plot.scSpectra.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra objects

```
sc.results <- screenSpectra(spectra)
summary(sc.results)</pre>
```

summaryPeaks

Description

This function generates a numerical summary of a collection of MassPeaks objects.

Usage

```
summaryPeaks(x, digits = 4)
```

Arguments

Х	A list of MassPeaks objects.
digits	Integer indicating the number of decimal places to be used.

Details

For each MassPeaks on the list this function provides summary statistics of m/z points, peak intensities and SNR thresholds (number, minimum, mean, standard deviation, median, mean absolute deviation, maximum).

Value

A data.frame containing summary information of a collection of MassPeaks objects.

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
data(type) # metadata
# Some pre-processing
sc.results <- screenSpectra(spectra, meta = type)
spectra <- sc.results$fspectra
type <- sc.results$fmeta
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
names(peaks) <- type$SpectID # spectra IDs are lost after removeBaseline()
# Summary of peak profile features (results for positions 10 to 20)
```

```
summaryPeaks(peaks[10:20])
```

summarySpectra Summary of mass spectra

Description

This function generates a numerical summary of a collection of MassSpectrum objects.

Usage

summarySpectra(x, digits = 4)

Arguments

х	A list of MassSpectrum objects.
digits	Integer indicating the number of decimal places to be used.

Details

For each MassSpectrum on the list this function provides summary statistics of m/z points and signal intensities (number, minimum, mean, standard deviation, median, mean absolute deviation, maximum).

Value

A data.frame containing summary information of a collection of MassSpectrum objects.

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
data(type) # metadata
# Summary of spectra features (results for 20 first mass spectra)
summarySpectra(spectra[1:20])
# Some pre-processing
sc.results <- screenSpectra(spectra, meta = type)
spectra <- sc.results$fspectra
type <- sc.results$fspectra
type <- sc.results$fmeta
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)</pre>
```

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transfIntensity

```
spectra <- removeBaseline(spectra)
names(spectra) <- type$SpectID # spectra IDs are lost with removeBaseline()
# Summary of spectra features (results for positions 10 to 20)
summarySpectra(spectra[10:20])</pre>
```

transfIntensity Transform intensity of MassSpectrum objects

Description

This function applies user-defined transformations on the intensities of MassSpectrum objects.

Usage

```
transfIntensity(x, fun = NULL, ...)
```

Arguments

х	A list of MassSpectrum objects.
fun	Name of an user-defined transformation function or any other pre-defined one in R.
	Other arguments.

Details

This function allows the user to define any sensible function to be applied on signal intensities. For logarithm and square root transformations it is equivalent to transformIntensity in the MALDIquant package.

Value

A list of MassSpectrum objects with signal intensities transformed according to fun.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects

Scale intensities into [0, 1] by dividing by their maximum value

scale.max <- function(x){x/max(x)} # define scaling function</pre>

scaled.spectra <- transfIntensity(spectra, fun = scale.max)</pre>

Compute natural logarithm of intensity values (using the pre-defined sqrt R function)

log.spectra <- transfIntensity(spectra, sqrt)</pre>

type

Example mass spectra metadata

Description

Metadata associated to the spectra data set containing information about isolate, biological and technical replicate numbers and mass spectra IDs.

Usage

data(type)

Format

The format is:

Isolate: Factor w/ 14 levels "280", "43", "45", ..: 2 2 2 2 2 2 2 2 2 2 ...

BioRep : int 1 1 1 1 2 2 2 2 3 3 ...

TechRep: int 1 2 3 4 1 2 3 4 1 2 ...

SpectID: Factor w/ 315 levels "160408C13","160408C14",..: 1 2 3 4 5 6 7 8 9 10 ...

Examples

data(type)
str(type)

wavSmoothing

Discrete wavelet transformation for MassSpectrum objects

Description

This function performs undecimated wavelet transform (UDWT) on mass spectra in MassSpectrum format. Alternatively, smoothing methods included in the MALDIquant package can be called.

Usage

writeIntensity

Arguments

х	A list of MassSpectrum objects.
method	Smoothing method used.
n.levels	Depth of the decomposition for wavelet-based smoothing.
	Other arguments.

Details

Note that from version 1.1.0 of MALDIrppa wavelet smoothing is conducted by maximal overlap discrete wavelet transformation and universal thresholding of coefficients based on methods available on the waveslim package. The optimal level of smoothing is determined by model-driven estimates of the thresholds. The parameter n.levels (values > 0 and <= log(length(x),2)) can be used to tweak the levels to obtain a smoother or rougher result.

Alternatively, smoothing methods SavitzkyGolay and MovingAverage from the MALDIquant package can be called directly from this function.

If the previous implementation of the wavelet method is required please download and install manually source files of version 1.0.5-1 from the archive of old sources of the package (https://CRAN.R-project.org/package=MALDIrppa).

Value

A list of MassSpectrum objects with denoised signal intensities.

Examples

```
# Load example data
```

data(spectra) # list of MassSpectra class objects

sqrt transformation and signal smoothing using UDWT

```
spectra <- transfIntensity(spectra, fun = "sqrt")
spectra <- wavSmoothing(spectra)</pre>
```

writeIntensity Write intensity matrix in different formats

Description

This function writes an intensity matrix as generated by intensityMatrix into a file in the R, csv, NEXUS or FASTA formats. For NEXUS format it allows to specify weights for peaks.

Usage

Arguments

x	Intensity matrix as obtained from intensityMatrix.
filename	A character string specifying a name for the destination file (filename extension not required).
format	One of R (default .RData file), text (comma-separated .csv file), NEXUS (.nex file) or FASTA (.fas file).
binary	Logical value. If TRUE, a binary version (1: peak presence, 0: peak absence) of x is saved (default FALSE).
labels	Optional vector of ID labels for the samples.
weights	Optional numeric vector of peak weights (NEXUS format).
	Additional arguments.

Details

This is a wrapper function to simplify the writing of an intensity matrix in different formats while adding some extra features. It includes the common NEXUS and FASTA formats as an extension of functions in the ape package to handle peak intensity data. It also allows for taxa/sample precomputed peak weights to be included in the NEXUS file. It checks whether the names meet NEXUS name conventions and gives them adequate format if not. A binary intensity matrix is always internally generated (binary = TRUE) when either the NEXUS or FASTA format is chosen. If any, NA values in x are assumed to denote zero intensity/peak absence and are then converted into zeros.

Value

No return value, file in selected format created on destination folder.

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
# Some pre-processing
spectra <- screenSpectra(spectra)$fspectra
spectra <- transformIntensity(spectra, method = "sqrt")
spectra <- wavSmoothing(spectra)
spectra <- removeBaseline(spectra)
peaks <- detectPeaks(spectra)
peaks <- alignPeaks(peaks, minFreq = 0.8)
# Intensity matrix
int <- intensityMatrix(peaks)
# Save as R file (saved to a temporary location as an example)
```

writeMetadata

writeMetadata Write metadata in different forma	ıts
---	-----

Description

This function is simply a wrapper to write the metadata associated with a collection of mass spectra into a file in either the R or csv format.

Usage

```
writeMetadata(x, filename = "Metadata", format = c("R", "csv"), ...)
```

Arguments

Х	Metadata in any sensible data format, preferably matrix or data.frame.
filename	A character string specifying a name for the destination file (filename extension not required).
format	One of R (default .RData file) or text (comma-separated .csv file).
	Other arguments.

Details

It uses either save or write.table to store the metadata. Check these functions for adequate data formats.

Value

No return value, file in selected format created on destination folder.

Examples

```
# Load example data
data(spectra) # list of MassSpectra class objects
data(type) # metadata
# Some pre-processing
sc.spectra <- screenSpectra(spectra, meta = type)</pre>
```

```
spectra <- sc.spectra$fspectra # filtered spectra</pre>
type <- sc.spectra$fmeta # filtered metadata</pre>
spectra <- transformIntensity(spectra, method = "sqrt")</pre>
spectra <- wavSmoothing(spectra)</pre>
spectra <- removeBaseline(spectra)</pre>
peaks <- detectPeaks(spectra)</pre>
peaks <- alignPeaks(peaks, minFreq = 0.8)</pre>
# Intensity matrix
int <- intensityMatrix(peaks)</pre>
# Save resulting data in R format (to a temporary location as an example)
writeIntensity(int, filename = file.path(tempdir(),"MyintMatrix"))
writeMetadata(type, filename = file.path(tempdir(),"MyMetadata"))
# Save resulting data in csv format (to a temporary location as an example)
writeIntensity(int, filename = file.path(tempdir(), "MyintMatrix"),
                format = "csv")
writeMetadata(type, filename = file.path(tempdir(),"MyMetadata"),
               format = "csv")
```

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