Package 'xcms'

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Title LC/MS and GC/MS Data Analysis

Author Colin A. Smith <csmith@scripps.edu>, Ralf Tautenhahn <rtautenh@gmail.com>, Steffen Neumann <sneumann@ipb-halle.de>, Paul Benton <hpbenton@scripps.edu>, Christopher Conley <cjconley@ucdavis.edu>, Johannes Rainer <Johannes.Rainer@eurac.edu>

Maintainer Steffen Neumann <sneumann@ipb-halle.de>

- **Depends** R (>= 2.14.0), methods, mzR (>= 1.1.6), BiocGenerics, ProtGenerics, Biobase
- Imports lattice, RColorBrewer
- Suggests faahKO, msdata, ncdf4, multtest, rgl, MassSpecWavelet (>= 1.5.2), RANN, RUnit, parallel
- Enhances Rgraphviz, Rmpi, XML
- **Description** Framework for processing and visualization of chromatographically separated and single-spectra mass spectral data. Imports from AIA/ANDI NetCDF, mzXML, mzData and mzML files. Preprocesses data for high-throughput, untargeted analyte profiling.

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URL http://metlin.scripps.edu/download/ and

https://github.com/sneumann/xcms

BugReports https://github.com/sneumann/xcms/issues/new

biocViews MassSpectrometry, Metabolomics

NeedsCompilation yes

R topics documented:

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absent-methods

Description

Determine which peaks are absent / present in a sample class

Arguments

object	xcmsSet-class object
class	Name of a sample class from sampclass
minfrac	minimum fraction of samples necessary in the class to be absent/present

Details

Determine which peaks are absent / present in a sample class The functions treat peaks that are only present because of fillPeaks correctly, i.e. does not count them as present.

Value

An logical vector with the same length as nrow(groups(object)).

Methods

```
object = "xcmsSet" absent(object, ...) present(object, ...)
```

See Also

group diffreport

AutoLockMass-methods Automatic parameter for Lock mass fixing AutoLockMass ~~

Description

AutoLockMass - This function decides where the lock mass scans are in the xcmsRaw object. This is done by using the scan time differences.

Arguments

object An xcmsRaw-class object

Value

AutoLockMass A numeric vector of scan locations corresponding to lock Mass scans

c-methods

Methods

object = "xcmsRaw" signature(object = "xcmsRaw")

Author(s)

Paul Benton, <hpaul.benton08@imperial.ac.uk>

Examples

```
## Not run: library(xcms)
library(faahKO) ## These files do not have this problem to correct for but just for an example
cdfpath <- system.file("cdf", package = "faahKO")
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)
xr<-xcmsRaw(cdffiles[1])
xr
##Lets assume that the lockmass starts at 1 and is every 100 scans
lockMass<-xcms:::makeacqNum(xr, freq=100, start=1)
## these are equalvent
lockmass2<-AutoLockMass(xr)
all((lockmass == lockmass2) == TRUE)
ob<-stitch(xr, lockMass)
## End(Not run)</pre>
```

c-methods Combine xcmsSet objects

Description

Combines the samples and peaks from multiple xcmsSet objects into a single object. Group and retention time correction data are discarded. The profinfo list is set to be equal to the first object.

Arguments

xs1	xcmsSet object
	xcmsSet objects

Value

A xcmsSet object.

Methods

xs1 = "xcmsRaw" c(xs1, ...)

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

xcmsSet-class

calibrate-methods Calibrate peaks for correcting unprecise m/z values

Description

Calibrate peaks of a xcmsSet via a set of known masses

Arguments

object	a xcmsSet object with uncalibrated mz
calibrants	a vector or a list of vectors with reference m/z-values
method	the used calibrating-method, see below
mzppm	the relative error used for matching peaks in ppm (parts per million)
mzabs	the absolute error used for matching peaks in Da
neighbours	the number of neighbours from wich the one with the highest intensity is used (instead of the nearest)
plotres	can be set to TRUE if wanted a result-plot showing the found m/z with the distances and the regression

Value

object	a xcmsSet with one ore more samples
calibrants	for each sample different calibrants can be used, if a list of m/z-vectors is given. The length of the list must be the same as the number of samples, alternatively a single vector of masses can be given which is used for all samples.
method	"shift" for shifting each m/z, "linear" does a linear regression and adds a linear term to each m/z. "edgeshift" does a linear regression within the range of the mz-calibrants and a shift outside.

Methods

object = "xcmsSet" ca	alibrate(object,	calibrants,method="1	linear",	mzabs=0.0001,	mzppm=5,	neigł
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See Also

xcmsSet-class,

collect-methods Collect MS^n peaks into xcmsFragments

Description

Collecting Peaks into xcmsFragmentss from several MS-runs using xcmsSet and xcmsRaw.

Arguments

object	(empty) xcmsFragments-class object
xs	A xcmsSet-class object which contains picked ms1-peaks from several exper- iments
compMethod	("floor", "round", "none"): compare-method which is used to find the parent peak of a MSnpeak through comparing the MZ-values of the MS1peaks with the MSnParentPeaks.
snthresh, mzga	
	these are the parameters for the getspec-peakpicker included in xcmsRaw.

Details

After running collect(xFragments,xSet) The peak table of the xcmsFragments includes the ms1Peaks from all experiments stored in a xcmsSet-object. Further it contains the relevant msN-peaks from the xcmsRaw-objects, which were created temporarily with the paths in xcmsSet.

Value

A matrix with columns:

peakID	unique identifier of every peak	
MSnParentPeakI)	
	PeakID of the parent peak of a msLevel>1 - peak, it is 0 if the peak is msLevel	
	1.	
msLevel	The msLevel of the peak.	
rt	retention time of the peak midpoint	
mz	the mz-Value of the peak	
intensity	the intensity of the peak	
sample	the number of the sample from the xcmsSet	
GroupPeakMSn	Used for grouped xcmsSet groups	
CollisionEnergy		
	The collision energy of the fragment	

Methods

object = "xcmsFragments" collect(object, ...)

diffreport-methods Create report of analyte differences

Description

Create a report showing the most significant differences between two sets of samples. Optionally create extracted ion chromatograms for the most significant differences.

Arguments

object	the xcmsSet object
class1	character vector with the first set of sample classes to be compared
class2	character vector with the second set of sample classes to be compared
filebase	base file name to save report, .tsv file and _eic will be appended to this name for the tabular report and EIC directory, respectively. if blank nothing will be saved
eicmax	number of the most significantly different analytes to create EICs for
eicwidth	width (in seconds) of EICs produced
sortpval	logical indicating whether the reports should be sorted by p-value
classeic	character vector with the sample classes to include in the EICs
value	<pre>intensity values to be used for the diffreport. If value="into", integrated peak intensities are used. If value="maxo", maximum peak intensities are used. If value="intb", baseline corrected integrated peak intensities are used (only available if peak detection was done by findPeaks.centWave).</pre>
metlin	mass uncertainty to use for generating link to Metlin metabolite database. the sign of the uncertainty indicates negative or positive mode data for M+H or M-H calculation. a value of FALSE or 0 removes the column
h	Numeric variable for the height of the eic and boxplots that are printed out.
W	Numeric variable for the width of the eic and boxplots print out made.
mzdec	Number of decimal places of title m/z values in the eic plot.
	optional arguments to be passed to mt.teststat

Details

This method handles creation of summary reports with statistics about which analytes were most significantly different between two sets of samples. It computes Welch's two-sample t-statistic for each analyte and ranks them by p-value. It returns a summary report that can optionally be written out to a tab-separated file.

Additionally, it does all the heavy lifting involved in creating superimposed extracted ion chromatograms for a given number of analytes. It does so by reading the raw data files associated with the samples of interest one at a time. As it does so, it prints the name of the sample it is currently reading. Depending on the number and size of the samples, this process can take a long time.

diffreport-methods

If a base file name is provided, the report (see Value section) will be saved to a tab separated file. If EICs are generated, they will be saved as 640x480 PNG files in a newly created subdirectory. However this parameter can be changed with the commands arguments. The numbered file names correspond to the rows in the report.

Chromatographic traces in the EICs are colored and labeled by their sample class. Sample classes take their color from the current palette. The color a sample class is assigned is dependent its order in the xcmsSet object, not the order given in the class arguments. Thus levels(sampclass(object))[1] would use color palette()[1] and so on. In that way, sample classes maintain the same color across any number of different generated reports.

When there are multiple sample classes, xcms will produce boxplots of the different classes and will generate a single anova p-value statistic. Like the eic's the plot number corresponds to the row number in the report.

Value

A data frame with the following columns:

fold	mean fold change (always greater than 1, see tstat for which set of sample classes was higher)
tstat	Welch's two sample t-statistic, positive for analytes having greater intensity in class2, negative for analytes having greater intensity in class1
pvalue	p-value of t-statistic
anova	p-value of the anova statistic if there are multiple classes
mzmed	median m/z of peaks in the group
mzmin	minimum m/z of peaks in the group
mzmax	maximum m/z of peaks in the group
rtmed	median retention time of peaks in the group
rtmin	minimum retention time of peaks in the group
rtmax	maximum retention time of peaks in the group
npeaks	number of peaks assigned to the group
Sample Classes	number samples from each sample class represented in the group
metlin	A URL to metlin for that mass
	one column for every sample class
Sample Names	integrated intensity value for every sample
	one column for every sample

Methods

```
object = "xcmsSet" diffreport(object, class1 = levels(sampclass(object))[1], class2 = levels(sampclass(object))[1],
```

See Also

xcmsSet-class, mt.teststat, palette

Description

A general function for asymmetric chromatographic peaks.

Usage

etg(x, H, t1, tt, k1, kt, lambda1, lambdat, alpha, beta)

Arguments

x	times to evaluate function at
Н	peak height
t1	time of leading edge inflection point
tt	time of trailing edge inflection point
k1	leading edge parameter
kt	trailing edge parameter
lambda1	leading edge parameter
lambdat	trailing edge parameter
alpha	leading edge parameter
beta	trailing edge parameter

Value

The function evaluated at times x.

Author(s)

Colin A. Smith, <csmith@scripps.edu>

References

Jianwei Li. Development and Evaluation of Flexible Empirical Peak Functions for Processing Chromatographic Peaks. Anal. Chem., 69 (21), 4452-4462, 1997. http://dx.doi.org/10.1021/ ac970481d

etg

Description

For each sample, identify peak groups where that sample is not represented. For each of those peak groups, integrate the signal in the region of that peak group and create a new peak.

Arguments

object	the xcmsSet object
method	the filling method

Details

After peak grouping, there will always be peak groups that do not include peaks from every sample. This method produces intensity values for those missing samples by integrating raw data in peak group region. According to the type of raw-data there are 2 different methods available. for filling gcms/lcms data the method "chrom" integrates raw-data in the chromatographic domain, whereas "MSW" is used for peaklists without retention-time information like those from direct-infusion spectra.

Value

A xcmsSet objects with filled in peak groups.

Methods

```
object = "xcmsSet" fillPeaks(object, method="")
```

See Also

xcmsSet-class, getPeaks

fillPeaks.chrom-methods

Integrate areas of missing peaks

Description

For each sample, identify peak groups where that sample is not represented. For each of those peak groups, integrate the signal in the region of that peak group and create a new peak.

Arguments

object	the xcmsSet object
nSlaves	number of slaves/cores to be used for parallel peak filling. MPI is used if in- stalled, otherwise the snow package is employed for multicore support. If none of the two packages is available it uses the parallel package for parallel process- ing on multiple CPUs of the current machine.
expand.mz	Expansion factor for the m/z range used for integration.
expand.rt	Expansion factor for the rentention time range used for integration.

Details

After peak grouping, there will always be peak groups that do not include peaks from every sample. This method produces intensity values for those missing samples by integrating raw data in peak group region. In a given group, the start and ending retention time points for integration are defined by the median start and end points of the other detected peaks. The start and end m/z values are similarly determined. Intensities can be still be zero, which is a rather unusual intensity for a peak. This is the case if e.g. the raw data was thresholded, and the integration area contains no actual raw intensities, or if one sample is miscalibrated, such thet the raw data points are (just) outside the integration area.

Importantly, if retention time correction data is available, the alignment information is used to more precisely integrate the propper region of the raw data. If the corrected retention time is beyond the end of the raw data, the value will be not-a-number (NaN).

Value

A xcmsSet objects with filled in peak groups (into and maxo).

Methods

object = "xcmsSet" fillPeaks.chrom(object, nSlaves=0,expand.mz=1,expand.rt=1)

See Also

xcmsSet-class, getPeaks fillPeaks

fillPeaks.MSW-methods Integrate areas of missing peaks in FTICR-MS data

Description

For each sample, identify peak groups where that sample is not represented. For each of those peak groups, integrate the signal in the region of that peak group and create a new peak.

Arguments

object the xcmsSet object

findMZ

Details

After peak grouping, there will always be peak groups that do not include peaks from every sample. This method produces intensity values for those missing samples by integrating raw data in peak group region. In a given group, the start and ending m/z values for integration are defined by the median start and end points of the other detected peaks.

Value

A xcmsSet objects with filled in peak groups.

Methods

object = "xcmsSet" fillPeaks.MSW(object)

See Also

xcmsSet-class, getPeaks fillPeaks

findMZ

Find fragment ions in xcmsFragment objects

Description

This is a method to find a fragment mass with a ppm window in a xcmsFragment object

Usage

findMZ(object, find, ppmE=25, print=TRUE)

Arguments

object	xcmsFragment object type
find	The fragment ion to be found
ppmE	the ppm error window for searching
print	If we should print a nice little report

Details

The method simply searches for a given fragment ion in an xcmsFragment object type given a certain ppm error window

Value

A data frame with the following columns:

PrecursorMz	The precursor m/z of the fragment
MSnParentPeakI)
	An index ID of the location of the precursor peak in the xcmsFragment object
msLevel	The level of the found fragment ion
rt	the Retention time of the found ion
mz	the actual m/z of the found fragment ion
intensity	The intensity of the fragment ion
sample	Which sample the fragment ion came from
GroupPeakMSn	an ID if the peaks were grouped by an xcmsSet grouping
CollisionEnergy	
	The collision energy of the precursor scan

Author(s)

H. Paul Benton, <hpaul.beonton08@imperial.ac.uk>

References

H. Paul Benton, D.M. Wong, S.A.Strauger, G. Siuzdak "XCMS2" Analytical Chemistry 2008

See Also

findneutral,

Examples

```
## Not run:
library(msdata)
mzdatapath <- system.file("iontrap", package = "msdata")
mzdatafiles<-list.files(mzdatapath, pattern = "extracted.mzData", recursive = TRUE, full.names = TRUE)
xs <- xcmsSet(mzdatafiles, method = "MS1")
##takes only one file from the file set
xfrag <- xcmsFragments(xs)
found<-findMZ(xfrag, 657.3433, 50)</pre>
```

End(Not run)

findneutral

Description

This is a method to find a neutral loss with a ppm window in a xcmsFragment object

Usage

findneutral(object, find, ppmE=25, print=TRUE)

Arguments

object	xcmsFragment object type
find	The neutral loss to be found
ppmE	the ppm error window for searching
print	If we should print a nice little report

Details

The method searches for a given neutral loss in an xcmsFragment object type given a certain ppm error window. The neutral losses are generated between neighbouring ions. The resulting data frame shows the whole scan in which the neutral loss was found.

Value

A data frame with the following columns:

PrecursorMz	The precursor m/z of the neutral losses
MSnParentPeakID	
	An index ID of the location of the precursor peak in the xcmsFragment object
msLevel	The level of the found fragment ion
rt	the Retention time of the found ion
mz	the actual m/z of the found fragment ion
intensity	The intensity of the fragment ion
sample	Which sample the fragment ion came from
GroupPeakMSn	an ID if the peaks were grouped by an xcmsSet grouping
CollisionEnerg	у
	The collision energy of the precursor scan

Author(s)

H. Paul Benton, <hpbenton@scripps.edu>

References

H. Paul Benton, D.M. Wong, S.A.Strauger, G. Siuzdak "XCMS²" Analytical Chemistry 2008

See Also

findMZ,

Examples

```
## Not run:
library(msdata)
mzdatapath <- system.file("iontrap", package = "msdata")
mzdatafiles<-list.files(mzdatapath, pattern = "extracted.mzData", recursive = TRUE, full.names = TRUE)
xs <- xcmsSet(mzdatafiles, method = "MS1")
##takes only one file from the file set
xfrag <- xcmsFragments(xs)
found<-findneutral(xfrag, 58.1455, 50)</pre>
```

End(Not run)

findPeaks-methods Feature detection for GC/MS and LC/MS Data - methods

Description

A number of peak pickers exist in XCMS. findPeaks is the generic method.

Arguments

object	xcmsRaw-class object
method	Method to use for peak detection. See details
	Optional arguments to be passed along

Details

Different algorithms can be used by specifying them with the method argument. For example to use the matched filter approach described by Smith et al (2006) one would use: findPeaks(object, method="matchedFilter" This is also the default.

Further arguments given by ... are passed through to the function implementing the method.

A character vector of *nicknames* for the algorithms available is returned by getOption("BioC")\$xcms\$findPeaks.methods If the nickname of a method is called "centWave", the help page for that specific method can be accessed with ?findPeaks.centWave.

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findPeaks.centWave-methods

Value

A matrix with columns:

mz	weighted (by intensity) mean of peak m/z across scans
mzmin	m/z of minimum step
mzmax	m/z of maximum step
rt	retention time of peak midpoint
rtmin	leading edge of peak retention time
rtmax	trailing edge of peak retention time
into	integrated area of original (raw) peak
maxo	maximum intensity of original (raw) peak
and additional columns depending on the choosen method.	

Methods

object = "xcmsRaw" findPeaks(object, ...)

See Also

findPeaks.matchedFilter findPeaks.centWave xcmsRaw-class

```
findPeaks.centWave-methods
```

Feature detection for high resolution LC/MS data

Description

Peak density and wavelet based feature detection for high resolution LC/MS data in centroid mode

Arguments

object	xcmsSet object
ppm	maxmial tolerated m/z deviation in consecutive scans, in ppm (parts per million)
peakwidth	Chromatographic peak width, given as range (min,max) in seconds
snthresh	signal to noise ratio cutoff, definition see below.
prefilter	prefilter=c(k,I). Prefilter step for the first phase. Mass traces are only re- tained if they contain at least k peaks with intensity >= I.
mzCenterFun	Function to calculate the m/z center of the feature: wMean intensity weighted mean of the feature m/z values, mean mean of the feature m/z values, apex use m/z value at peak apex, wMeanApex3 intensity weighted mean of the m/z value at peak apex and the m/z value left and right of it, meanApex3 mean of the m/z value at peak apex and the m/z value left and right of it.

integrate	Integration method. If =1 peak limits are found through descent on the mexican hat filtered data, if =2 the descent is done on the real data. Method 2 is very accurate but prone to noise, while method 1 is more robust to noise but less exact.
mzdiff	minimum difference in m/z for peaks with overlapping retention times, can be negative to allow overlap
fitgauss	logical, if TRUE a Gaussian is fitted to each peak
scanrange	scan range to process
noise	optional argument which is useful for data that was centroided without any inten- sity threshold, centroids with intensity < noise are omitted from ROI detection
sleep	number of seconds to pause between plotting peak finding cycles
verbose.columns	
	logical, if TRUE additional peak meta data columns are returned
ROI.list	A optional list of ROIs that represents detected mass traces (ROIs). If this list is empty (default) then centWave detects the mass trace ROIs, otherwise this step is skipped and the supplied ROIs are used in the peak detection phase. Each ROI object in the list has the following slots: scmin start scan index, scmax end scan index, mzmin minimum m/z, mzmax maximum m/z, length number of scans, intensity summed intensity.

Details

This algorithm is most suitable for high resolution LC/{TOF,OrbiTrap,FTICR}-MS data in centroid mode. In the first phase of the method mass traces (characterised as regions with less than ppm m/z deviation in consecutive scans) in the LC/MS map are located. In the second phase these mass traces are further analysed. Continuous wavelet transform (CWT) is used to locate chromatographic peaks on different scales.

Value

A matrix with columns:

mz	weighted (by intensity) mean of peak m/z across scans
mzmin	m/z peak minimum
mzmax	m/z peak maximum
rt	retention time of peak midpoint
rtmin	leading edge of peak retention time
rtmax	trailing edge of peak retention time
into	integrated peak intensity
intb	baseline corrected integrated peak intensity
maxo	maximum peak intensity
sn	Signal/Noise ratio, defined as (maxo - baseline)/sd, where maxo is the maximum peak intensity, baseline the estimated baseline value and sd the standard deviation of local chromatographic noise.

findPeaks.massifquant-methods

egauss	RMSE of Gaussian fit
	if verbose.columns is TRUE additionally :
mu	Gaussian parameter mu
sigma	Gaussian parameter sigma
h	Gaussian parameter h
f	Region number of m/z ROI where the peak was localised
dppm	m/z deviation of mass trace across scans in ppm
scale	Scale on which the peak was localised
scpos	Peak position found by wavelet analysis
scmin	Left peak limit found by wavelet analysis (scan number)
scmax	Right peak limit found by wavelet analysis (scan number)

Methods

object = ''xcmsRaw''	findPeaks.	<pre>centWave(object,</pre>	ppm=25,	<pre>peakwidth=c(20,50),</pre>	snthresh=10,	prefilt
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Author(s)

Ralf Tautenhahn

References

Ralf Tautenhahn, Christoph B\"ottcher, and Steffen Neumann "Highly sensitive feature detection for high resolution LC/MS" BMC Bioinformatics 2008, 9:504

See Also

findPeaks-methods xcmsRaw-class

Description

Massifquant is a Kalman filter (KF) based feature detection for XC-MS data in centroid mode (currently in experimental stage). Optionally allows for calling the method "centWave" on features discovered by Massifquant to further refine the feature detection; to do so, supply any additional parameters specific to centWave (even more experimental). The method may be conveniently called through the xcmsSet(...) method.

Arguments	
	The following arguments are specific to Massifquant. Any additional arguments supplied must correspond as specified by the method findPeaks.centWave.
	An xcmsRaw object.
objėċtalValue	Numeric: Suggested values: (0.1-3.0). This setting helps determine the the Kalman Filter prediciton margin of error. A real centroid belonging to a bonafide feature must fall within the KF prediction margin of error. Much like in the construction of a confidence interval, criticalVal loosely translates to be a multiplier of the standard error of the prediction reported by the Kalman Filter. If the features in the XC-MS sample have a small mass deviance in ppm error, a smaller critical value might be better and vice versa.
consecMissedLin	
	Integer: Suggested values:(1,2,3). While a feature is in the proces of being detected by a Kalman Filter, the Kalman Filter may not find a predicted centroid in every scan. After 1 or more consecutive failed predictions, this setting informs Massifquant when to stop a Kalman Filter from following a candidate feature.
prefilter	Numeric Vector: (Positive Integer, Positive Numeric): The first argument is only used if (withWave = 1); see centWave for details. The second argument specifies the minimum threshold for the maximum intensity of a feature that must be met.
peakwidth	Integer Vector: (Positive Integer, Positive Integer): Only the first argument is used for Massifquant, which specifices the minimum feature length in time scans. If centWave is used, then the second argument is the maximum feature length subject to being greater than the minimum feature length.
ppm	The minimum estimated parts per million mass resolution a feature must possess.
unions	Integer: set to 1 if apply t-test union on segmentation; set to 0 if no t-test to be applied on chromatographically continous features sharing same m/z range. Ex- planation: With very few data points, sometimes a Kalman Filter stops tracking a feature prematurely. Another Kalman Filter is instantiated and begins follow- ing the rest of the signal. Because tracking is done backwards to forwards, this algorithmic defect leaves a real feature divided into two segments or more. With this option turned on, the program identifies segmented features and combines them (merges them) into one with a two sample t-test. The potential danger of this option is that some truly distinct features may be merged.
withWave	Integer: set to 1 if turned on; set to 0 if turned off. Allows the user to find features first with Massifquant and then filter those features with the second phase of centWave, which includes wavelet estimation.
checkBack	Integer: set to 1 if turned on; set to 0 if turned off. The convergence of a Kalman Filter to a feature's precise m/z mapping is very fast, but sometimes it incorporates erroneous centroids as part of a feature (especially early on). The "scan-Back" option is an attempt to remove the occasional outlier that lies beyond the converged bounds of the Kalman Filter. The option does not directly affect identification of a feature because it is a postprocessing measure; it has not shown to be a extremely useful thus far and the default is set to being turned off.

Details

This algorithm's performance has been tested rigorously on high resolution LC/{OrbiTrap, TOF}-MS data in centroid mode. Simultaneous kalman filters identify features and calculate their area under the curve. The default parameters are set to operate on a complex LC-MS Orbitrap sample. Users will find it useful to do some simple exploratory data analysis to find out where to set a minimum intensity, and identify how many scans an average feature spans. The "consecMissedLimit" parameter has yielded good performance on Orbitrap data when set to (2) and on TOF data it was found best to be at (1). This may change as the algorithm has yet to be tested on many samples. The "criticalValue" parameter is perhaps most dificult to dial in appropriately and visual inspection of peak identification is the best suggested tool for quick optimization. The "ppm" and "checkBack" parameters have shown less influence than the other parameters and exist to give users flexibility and better accuracy.

Value

If the method findPeaks.massifquant(...) is used, then a matrix is returned with rows corresponding to features, and properties of the features listed with the following column names. Otherwise, if centWave feature is used also (withWave = 1), or Massifquant is called through the xcmsSet(...) method, then their corresponding return values are used.

mz	weighted m/z mean (weighted by intensity) of the feature
mzmin	m/z lower boundary of the feature
mzmax	m/z upper boundary of the feature
rtmin	starting scan time of the feature
rtmax	starting scan time of the feature
into	the raw quantitation (area under the curve) of the feature.
area	feature area that is not normalized by the scan rate.

Methods

object = "xcmsRaw" findPeaks.massifquant(object, ppm=10, peakwidth=c(20,50), snthresh=10, prefil

Author(s)

Christopher Conley

References

Submitted for review. Christopher Conley, Ralf J .O Torgrip. Ryan Taylor, and John T. Prince. "Massifquant: open-source Kalman filter based XC-MS feature detection". August 2013.

See Also

findPeaks-methods xcmsSet xcmsRaw xcmsRaw-class

Examples

findPeaks.matchedFilter-methods

Feature detection in the chromatographic time domain

Description

Find peaks in extracted the chromatographic time domain of the profile matrix.

Arguments

object	xcmsRaw object
fwhm	full width at half maximum of matched filtration gaussian model peak. Only used to calculate the actual sigma, see below.
sigma	standard deviation (width) of matched filtration model peak
max	maximum number of peaks per extracted ion chromatogram
snthresh	signal to noise ratio cutoff
step	step size to use for profile generation
steps	number of steps to merge prior to filtration
mzdiff	minimum difference in m/z for peaks with overlapping retention times
index	return indicies instead of values for m/z and retention times
sleep	number of seconds to pause between plotting peak finding cycles
scanrange	scan range to process

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Value

A matrix with columns:

mz	weighted (by intensity) mean of peak m/z across scans
mzmin	m/z of minimum step
mzmax	m/z of maximum step
rt	retention time of peak midpoint
rtmin	leading edge of peak retention time
rtmax	trailing edge of peak retention time
into	integrated area of original (raw) peak
intf	integrated area of filtered peak
maxo	maximum intensity of original (raw) peak
maxf	maximum intensity of filtered peak
i	rank of peak identified in merged EIC (<= max)
sn	signal to noise ratio of the peak

Methods

object = ''xcmsRaw''	<pre>findPeaks.matchedFilter(object</pre>	, fwhm = 30,	, sigma = fwhm/2.3548,	max = 5,	snt
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Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

findPeaks-methods xcmsRaw-class

findPeaks.MS1-methods Collecting MS1 precursor peaks

Description

Collecting Tandem MS or MS\$^n\$ Mass Spectrometry precursor peaks as annotated in XML raw file

Arguments

object xcmsRaw object

Details

Some mass spectrometers can acquire MS1 and MS2 (or MS\$^n\$ scans) quasi simultaneously, e.g. in data dependent tandem MS or DDIT mode.

Since xcmsFragments attaches *all* MS\$^n\$ peaks to MS1 peaks in xcmsSet, it is important that findPeaks and xcmsSet do not miss any MS1 precursor peak.

To be sure that all MS1 precursor peaks are in an xcmsSet, findPeaks.MS1 does not do an actual peak picking, but simply uses the annotation stored in mzXML, mzData or mzML raw files.

This relies on the following XML tags:

```
mzData: <spectrum id="463"> <spectrumInstrument msLevel="2">
<cvParam cvLabel="psi" accession="PSI:1000039" name="TimeInSeconds" value="92.7743"/>
</spectrumInstrument> <precursor msLevel="1" spectrumRef="461">
<cvParam cvLabel="psi" accession="PSI:1000040" name="MassToChargeRatio" value="462.091"/>
<cvParam cvLabel="psi" accession="PSI:1000042" name="Intensity" value="366.674"/>
</precursor> </spectrum>
```

```
mzXML: <scan num="17" msLevel="2" retentionTime="PT1.5224S"> <precursorMz precursorIntensit
</scan>
```

Several mzXML and mzData converters are known to create incomplete files, either without intensities (they will be set to 0) or without the precursor retention time (then a reasonably close rt will be chosen. NYI).

Value

A matrix with columns:

mz,	mzmin,	mzmax
		annotated MS1 precursor selection mass
rt,	rtmin,	rtmax
		annotated MS1 precursor retention time
inte	o, maxo	sn annotated MS1 precursor intensity

Methods

object = "xcmsRaw" findPeaks.MS1(object)

Author(s)

Steffen Neumann, <sneumann@ipb-halle.de>

See Also

findPeaks-methods xcmsRaw-class

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findPeaks.MSW-methods Feature detection for single-spectrum non-chromatography MS data

Description

Processing Mass Spectrometry direct-injection spectrum by using wavelet based algorithm.

Arguments

object	xcmsSet object	
snthresh	signal to noise ratio cutoff	
scales	scales of CWT	
nearbyPeak	Determine whether to include the nearby small peaks of major peaks. TRUE by default	
sleep	number of seconds to pause between plotting peak finding cycles	
verbose.columns		
	additional peak meta data columns are returned	

Details

This is a wrapper around the peak picker in the bioconductor package MassSpecWavelet calling 'cwt', 'get.localMaximum.cwt', 'get.ridge', 'identify.majorPeaks' and tuneIn.peakInfo.

Value

A matrix with columns:

mz	m/z value of the peak at the centroid position
mzmin	m/z value at the start-point of the peak
mzmax	m/z value at the end-point of the peak
rt	always -1
rtmin	always -1
rtmax	always -1
into	integrated area of original (raw) peak
maxo	intensity of original (raw) peak at the centroid position
intf	always NA
maxf	maximum MSW-filter response of the peak
sn	Signal/Noise ratio

Methods

object = ''xcmsRaw''	findPeaks.MSW(object, snthresh=3,	<pre>scales=seq(1,22,3), nearbyPeak=TRUE,</pre>	р
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Author(s)

Steffen Neumann, Joachim kutzera, <sneumann|jkutzer@ipb-halle.de>

See Also

findPeaks-methods xcmsRaw-class peakDetectionCWT

getEIC-methods Get extracted ion chromatograms for specified m/z ranges

Description

Generate multiple extracted ion chromatograms for m/z values of interest. For xcmsSet objects, reread original raw data and apply precomputed retention time correction, if applicable.

Arguments

the xcmsRaw or xcmsSet object
either a two column matrix with minimum or maximum m/z or a matrix of any dimensions containing columns mzmin and mzmax. If not specified, the method for xcmsRaw returns the base peak chromatogram (BPC, i.e. the most intense signal for each RT across all m/z).
for xcmsSet objects, if left blank the group data will be used instead
a two column matrix the same size as mzrange with minimum and maximum retention times between which to return EIC data points. If not specified, the method returns the chromatogram for the full RT range.
for xcmsSet objects, it may also be a single number specifying the time window around the peak to return EIC data points
step size to use for profile generation
either character vector with names or integer vector with indicies of peak groups for which to get EICs
either character vector with names or integer vector with indicies of samples for which to get EICs
"corrected" for using corrected retention times, or "raw" for using raw reten- tion times

Value

For xcmsSet and xcmsRaw objects, an xcmsEIC object.

Methods

```
object = "xcmsRaw" getEIC(object, mzrange, rtrange = NULL, step = 0.1)
object = "xcmsSet" getEIC(object, mzrange, rtrange = 200, groupidx,
```

sampleidx = sampnames(c

getPeaks-methods

See Also

xcmsRaw-class, xcmsSet-class, xcmsEIC-class

getPeaks-methods Get peak intensities for specified regions

Description

Integrate extracted ion chromatograms in pre-defined defined regions. Return output similar to findPeaks.

Arguments

object	the xcmsSet object
peakrange	matrix or data frame with 4 columns: mzmin, mzmax, rtmin, rtmax (they must be in that order or named)
step	step size to use for profile generation

Value

A matrix with columns:

i	rank of peak identified in merged EIC (<= max), always NA
mz	weighted (by intensity) mean of peak m/z across scans
mzmin	m/z of minimum step
mzmax	m/z of maximum step
ret	retention time of peak midpoint
retmin	leading edge of peak retention time
retmax	trailing edge of peak retention time
into	integrated area of original (raw) peak
intf	integrated area of filtered peak, always NA
maxo	maximum intensity of original (raw) peak
maxf	maximum intensity of filtered peak, always NA

Methods

```
object = "xcmsRaw" getPeaks(object, peakrange, step = 0.1)
```

See Also

xcmsRaw-class

getScan-methods

Description

Return the data from a single mass scan using the numeric index of the scan as a reference.

Arguments

object	the xcmsRaw object
scan	integer index of scan. if negative, the index numbered from the end
mzrange	limit data points returned to those between in the range, range(mzrange)

Value

A matrix with two columns:

mz	m/z values
intensity	intensity values

Methods

object = "xcmsRaw" getScan(object, scan, mzrange = numeric()) getMsnScan(object, scan, mzrange = numer

See Also

xcmsRaw-class, getSpec

getSpec-methods *Get average m/z and intensity values for multiple mass scans*

Description

Return full-resolution averaged data from multiple mass scans.

Arguments

object	the xcmsRaw object
	arguments passed to profRange used to sepecify the spectral segments of inter- est for averaging

Details

Based on the mass points from the spectra selected, a master unique list of masses is generated. Every spectra is interpolated at those masses and then averaged.

Value

A matrix with two columns:

mz	m/z values
intensity	intensity values

Methods

object = "xcmsRaw" getSpec(object, ...)

See Also

xcmsRaw-class, profRange, getScan

getXcmsRaw-methods Load the raw data for one or more files in the xcmsSet

Description

Reads the raw data applies evential retention time corrections and waters Lock mass correction and returns it as an xcmsRaw object (or list of xcmsRaw objects) for one or more files of the xcmsSet object.

Arguments

object	the xcmsSet object
sampleidx	The index of the sample for which the raw data should be returned. Can be a single number or a numeric vector with the indices. Alternatively, the file name can be specified.
profmethod	The profile method.
profstep	The profile step.
rt	Whether corrected or raw retention times should be returned.
	Additional arguments submitted to the xcmsRaw function.

Value

A single xcmsRaw object or a list of xcmsRaw objects.

Methods

```
object = "xcmsSet" getXcmsRaw(object, sampleidx=1, profmethod=profinfo(object)$method, profstep=pro
)
```

Author(s)

Johannes Rainer, <johannes.rainer@eurac.edu>

See Also

xcmsRaw-class,

group-methods Gro

Group peaks from different samples together

Description

A number of grouping (or alignment) methods exist in XCMS. group is the generic method.

Arguments

object	xcmsSet-class object
method	Method to use for grouping. See details.
	Optional arguments to be passed along

Details

Different algorithms can be used by specifying them with the method argument. For example to use the density-based approach described by Smith et al (2006) one would use: group(object, method="density"). This is also the default.

Further arguments given by ... are passed through to the function implementing the method.

A character vector of *nicknames* for the algorithms available is returned by getOption("BioC")\$xcms\$group.methods. If the nickname of a method is called "mzClust", the help page for that specific method can be accessed with ?group.mzClust.

Value

An xcmsSet object with peak group assignments and statistics.

Methods

```
object = "xcmsSet" group(object, ...)
```

See Also

group.density group.mzClust group.nearest xcmsSet-class,

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group.density

Description

Group peaks together across samples using overlapping m/z bins and calculation of smoothed peak distributions in chromatographic time.

Arguments

object	the xcmsSet object
minfrac	minimum fraction of samples necessary in at least one of the sample groups for it to be a valid group
minsamp	minimum number of samples necessary in at least one of the sample groups for it to be a valid group
bw	bandwidth (standard deviation or half width at half maximum) of gaussian smooth- ing kernel to apply to the peak density chromatogram
mzwid	width of overlapping m/z slices to use for creating peak density chromatograms and grouping peaks across samples
max	maximum number of groups to identify in a single m/z slice
sleep	seconds to pause between plotting successive steps of the peak grouping algo- rithm. peaks are plotted as points showing relative intensity. identified groups are flanked by dotted vertical lines.

Value

An xcmsSet object with peak group assignments and statistics.

Methods

object = ''xcmsSet''	group(object, bw	= 30, minfrac	= 0.5, minsamp = 1,	mzwid = 0.25, max = 50, sle
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See Also

xcmsSet-class, density

group.mzClust

Description

Runs high resolution alignment on single spectra samples stored in a given xcmsSet.

Arguments

object	a xcmsSet with peaks
mzppm	the relative error used for clustering/grouping in ppm (parts per million)
mzabs	the absolute error used for clustering/grouping
minsamp	set the minimum number of samples in one bin
minfrac	set the minimum fraction of each class in one bin

Value

Returns a xcmsSet with slots groups and groupindex set.

Methods

object = "xcmsSet"	group(object, method="mzClust", mzppm = 20, mzabs = 0, minsamp = 1, minfrac=0))

References

Saira A. Kazmi, Samiran Ghosh, Dong-Guk Shin, Dennis W. Hill and David F. Grant *Alignment of high resolution mass spectra: development of a heuristic approach for metabolomics*. Metabolomics, Vol. 2, No. 2, 75-83 (2006)

See Also

xcmsSet-class,

Examples

Description

Group peaks together across samples by creating a master peak list and assigning corresponding peaks from all samples. It is inspired by the alignment algorithm of mzMine. For further details check http://mzmine.sourceforge.net/ and

Katajamaa M, Miettinen J, Oresic M: MZmine: Toolbox for processing and visualization of mass spectrometry based molecular profile data. Bioinformatics (Oxford, England) 2006, 22:634?636.

Currently, there is no equivalent to minfrac or minsamp.

Arguments

object	the xcmsSet object
mzVsRTbalance	Multiplicator for mz value before calculating the (euclidean) distance between two peaks.
mzCheck	Maximum tolerated distance for mz.
rtCheck	Maximum tolerated distance for RT.
knn	Number of nearest Neighbours to check

Value

An xcmsSet object with peak group assignments and statistics.

Methods

object = "xcmsSet" group(object, mzVsRTbalance=10, mzCheck=0.2, rtCheck=15, kNN=10)

See Also

xcmsSet-class, group.density and group.mzClust

Examples

```
## Not run: library(xcms)
library(faahKO) ## These files do not have this problem to correct for but just for an example
cdfpath <- system.file("cdf", package = "faahKO")
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)</pre>
```

```
xset<-xcmsSet(cdffiles)</pre>
```

```
gxset<-group(xset, method="nearest")
## this is the same as
# gxset<-group.nearest(xset)
nrow(gxset@groups) == 1096 ## the number of features before minFrac</pre>
```

```
post.minFrac<-function(object, minFrac=0.5){</pre>
ix.minFrac<-sapply(1:length(unique(sampclass(object))), function(x, object, mf){</pre>
meta<-groups(object)</pre>
minFrac.idx<-numeric(length=nrow(meta))</pre>
idx<-which(meta[,levels(sampclass(object))[x]] >= mf*length(which(levels(sampclass(object))[x] == sampclass(obj
minFrac.idx[idx]<-1</pre>
return(minFrac.idx)
}, object, minFrac)
ix.minFrac<-as.logical(apply(ix.minFrac, 1, sum))</pre>
ix<-which(ix.minFrac == TRUE)</pre>
return(ix)
}
## using the above function we can get a post processing minFrac
idx<-post.minFrac(gxset)</pre>
gxset.post<-gxset ## copy the xcmsSet object</pre>
gxset.post@groupidx<-gxset@groupidx[idx]
gxset.post@groups<-gxset@groups[idx,]</pre>
nrow(gxset.post@groups) == 465 ## this is the number of features after minFrac
## End(Not run)
```

groupnames-methods Generate unque names for peak groups

Description

Allow linking of peak group data between classes using unique group names that remain the same as long as no re-grouping occurs.

Arguments

object	the xcmsSet or xcmsEIC object
mzdec	number of decimal places to use for m/z
rtdec	number of decimal places to use for retention time
template	a character vector with existing group names whose format should be emulated

Value

A character vector with unique names for each peak group in the object. The format is M[m/z]T[time in seconds].

Methods

```
object = "xcmsSet" (object, mzdec = 0, rtdec = 0, template = NULL)
object = "xcmsEIC" (object)
```

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groupval-methods

See Also

xcmsSet-class, xcmsEIC-class

groupval-methods *Extract a matrix of peak values for each group*

Description

Generate a matrix of peak values with rows for every group and columns for every sample. The value included in the matrix can be any of the columns from the xcmsSet peaks slot matrix. Collisions where more than one peak from a single sample are in the same group get resolved with one of several user-selectable methods.

Arguments

object	the xcmsSet object
method	conflict resolution method, "medret" to use the peak closest to the median re- tention time or "maxint" to use the peak with the highest intensity
value	name of peak column to enter into returned matrix, or "index" for index to the corresponding row in the peaks slot matrix
intensity	if method == "maxint", name of peak column to use for intensity

Value

A matrix with with rows for every group and columns for every sample. Missing peaks have NA values.

Methods

object = "xcmsSet" groupval(object, method = c("medret", "maxint"), value = "index", inter value = "index", inter

See Also

xcmsSet-class

image-methods

Description

Create log intensity false-color image of a xcmsRaw object plotted with m/z and retention time axes

Arguments

х	xcmsRaw object
col	vector of colors to use for for the image
	arguments for profRange

Methods

```
x = "xcmsRaw" image(x, col = rainbow(256), ...)
```

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

xcmsRaw-class

levelplot-methods Plot log intensity image of a xcmsRaw object

Description

Create an image of the raw (profile) data m/z against retention time, with the intensity color coded.

Arguments

х	xcmsRaw object.
log	Whether the intensity should be log transformed.
col.regions	The color ramp that should be used for encoding of the intensity.
rt	wheter the original (rt="raw") or the corrected (rt="corrected") retention times should be used.
	Arguments for profRange.

Methods

x = "xcmsRaw"	<pre>levelplot(x, log=TRUE, col.regions=colorRampPalette(brewer.pal(9,</pre>	"YlOrRd"))(25
x = ''xcmsSet''	<pre>levelplot(x, log=TRUE, col.regions=colorRampPalette(brewer.pal(9,</pre>	"YlOrRd"))(256

Author(s)

Johannes Rainer, <johannes.rainer@eurac.edu>

See Also

xcmsRaw-class, xcmsSet-class

loadRaw-methods Read binary data from a source

Description

This function extracts the raw data which will be used an xcmsRaw object. Further processing of data is done in the xcmsRaw constructor.

Arguments

object Specification of a data source (such as a file name or database query)

Details

The implementing methods decide how to gather the data.

Value

A list containing elements describing the data source. The rt, scanindex, tic, and acquisitionNum components each have one entry per scan. They are "parallel" in the sense that rt[1], scanindex[1], and acquisitionNum[1] all refer to the same scan. The list containst he following components:

rt	Numeric vector with acquisition time (in seconds) for each scan
tic	Numeric vector with Total Ion Count for each scan
scanindex	Integer vector with starting positions of each scan in the mz and intensity components. It is an exclusive offset, so scanindex[i] is the offset in mz and intensity <i>before</i> the beginning of scan i. This means that the mz (respectively intensity) values for scan i would be from scanindex[i] + 1 to scanindex[i + 1]
mz	Concatenated vector of m/z values for all scans
intensity	Concatenated vector of intensity values for all scans

Methods

signature(object = "xcmsSource") Uses loadRaw, xcmsSource-method to extract raw data. Subclasses of xcmsSource can provide different ways of fetching data.

Author(s)

Daniel Hackney, <dan@haxney.org>

See Also

xcmsRaw-class, xcmsSource

medianFilter Apply a median filter to a matrix

Description

For each element in a matix, replace it with the median of the values around it.

Usage

medianFilter(x, mrad, nrad)

Arguments

х	numeric matrix to median filter
mrad	number of rows on either side of the value to use for median calculation
nrad	number of rows on either side of the value to use for median calculation

Value

A matrix whose values have been median filtered

Author(s)

Colin A. Smith, <csmith@scripps.edu>

Examples

```
mat <- matrix(1:25, nrow=5)
mat
medianFilter(mat, 1, 1)</pre>
```

msn2xcmsRaw

Description

The MS2 and MSn data is stored in separate slots, and can not directly be used by e.g. findPeaks(). msn2xcmsRaw() will copy the MSn spectra into the "normal" xcmsRaw slots.

Usage

msn2xcmsRaw(xmsn)

Arguments

xmsn

an object of class xcmsRaw that contains spectra read with includeMSn=TRUE

Details

The default gap value is determined from the 90th percentile of the pair-wise differences between adjacent mass values.

Value

An xcmsRaw object

Author(s)

Steffen Neumann <sneumann@ipb-halle.de>

See Also

xcmsRaw,

Examples

```
msnfile <- system.file("microtofq/MSMSpos20_6.mzML", package = "msdata")
xrmsn <- xcmsRaw(msnfile, includeMSn=TRUE)
xr <- msn2xcmsRaw(xrmsn)
p <- findPeaks(xr, method="centWave")</pre>
```

peakPlots-methods

Description

Plot extracted ion chromatograms for many peaks simultaneously, indicating peak integration start and end points with vertical grey lines.

Arguments

object	the xcmsRaw object
peaks	matrix with peak information as produced by findPeaks
figs	two-element vector describing the number of rows and the number of columns of peaks to plot, if missing then an approximately square grid that will fit the number of peaks supplied
width	width of chromatogram retention time to plot for each peak

Details

This function is intended to help graphically analyze the results of peak picking. It can help estimate the number of false positives and improper integration start and end points. Its output is very compact and tries to waste as little space as possible. Each plot is labeled with rounded m/z and retention time separated by a space.

Methods

```
signature(object = "xcmsSet") plotPeaks(object, peaks, figs, width = 200)
```

See Also

xcmsRaw-class, findPeaks, split.screen

peakTable-methods Create report of aligned peak intensities

Description

Create a report showing all aligned peaks.

object	the xcmsSet object
filebase	base file name to save report, .tsv file and _eic will be appended to this name for the tabular report and EIC directory, respectively. if blank nothing will be saved
	arguments passed down to groupval, which provides the actual intensities.

Details

This method handles creation of summary reports similar to diffreport. It returns a summary report that can optionally be written out to a tab-separated file.

If a base file name is provided, the report (see Value section) will be saved to a tab separated file.

Value

A data frame with the following columns:

mz	median m/z of peaks in the group
mzmin	minimum m/z of peaks in the group
mzmax	maximum m/z of peaks in the group
rt	median retention time of peaks in the group
rtmin	minimum retention time of peaks in the group
rtmax	maximum retention time of peaks in the group
npeaks	number of peaks assigned to the group
Sample Classes	number samples from each sample class represented in the group
	one column for every sample class
Sample Names	integrated intensity value for every sample
	one column for every sample

Methods

object = "xcmsSet" peakTable(object, filebase = character(), ...)

See Also

xcmsSet-class,

Examples

```
## Not run:
library(faahKO)
cdfpath <- system.file("cdf", package = "faahKO")
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)
xs<-xcmsSet(cdf files)
xs<-group(xs)
peakTable(xs, filebase="peakList")
```

End(Not run)

plot.xcmsEIC

Description

Batch plot a list of extracted ion chromatograms to the current graphics device.

Arguments

x	the xcmsEIC object
У	optional xcmsSet object with peak integration data
groupidx	either character vector with names or integer vector with indicies of peak groups for which to plot EICs
sampleidx	either character vector with names or integer vector with indicies of samples for which to plot EICs
rtrange	a two column matrix with minimum and maximum retention times between which to return EIC data points
	if it has the same number of rows as the number groups in the xcmsEIC object, then sampleidx is used to subset it. otherwise, it is repeated over the length of sampleidx
	it may also be a single number specifying the time window around the peak for which to plot EIC data
col	color to use for plotting extracted ion chromatograms. if missing and y is speci- fied, colors are taken from unclass(sampclass(y)) and the default palette if it is the same length as the number groups in the xcmsEIC object, then sampleidx is used to subset it. otherwise, it is repeated over the length of sampleidx
legtext	text to use for legend. if NULL and y is specified, legend text is taken from the sample class information found in the xcmsSet
peakint	logical, plot integrated peak area with darkened lines (requires that y also be specified)
sleep	seconds to pause between plotting EICs
	other graphical parameters

Value

A xcmsSet object.

Methods

```
x = "xcmsEIC" plot.xcmsEIC(x, y, groupidx = groupnames(x), sampleidx = sampnames(x), rtrange = x@rtr
```

Author(s)

Colin A. Smith, <csmith@scripps.edu>

plotChrom-methods

See Also

xcmsEIC-class, png, pdf, postscript,

plotChrom-methods *Plot extracted ion chromatograms from the profile matrix*

Description

Uses the pre-generated profile mode matrix to plot averaged or base peak extracted ion chromatograms over a specified mass range.

Arguments

object	the xcmsRaw object
base	logical, plot a base-peak chromatogram
ident	logical, use mouse to identify and label peaks
fitgauss	logical, fit a gaussian to the largest peak
vline	numeric vector with locations of vertical lines
	arguments passed to profRange

Value

If ident == TRUE, an integer vector with the indecies of the points that were identified. If fitgauss == TRUE, a nls model with the fitted gaussian. Otherwise a two-column matrix with the plotted points.

Methods

object = "xcmsRaw" plotChrom(object, base = FALSE, ident = FALSE,

fitgauss = FALSE, vlir

See Also

xcmsRaw-class

plotEIC-methods

Description

Plot extracted ion chromatogram for m/z values of interest. The raw data is used in contrast to plotChrom which uses data from the profile matrix.

Arguments

object	xcmsRaw object
mzrange	m/z range for EIC. Uses the full m/z range by default.
rtrange	retention time range for EIC. Uses the full retention time range by default.
scanrange	scan range for EIC
mzdec	Number of decimal places of title m/z values in the eic plot.
type	Speficies how the data should be plotted (by default as a line).
add	If the EIC should be added to an existing plot.
	Additional parameters passed to the plotting function (e.g. col etc).

Value

A two-column matrix with the plotted points.

Methods

object = "xcmsRaw"	plotEIC(object, m	<pre>mzrange = numeric(),</pre>	<pre>rtrange = numeric(),</pre>	scanrange = numeri
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Author(s)

Ralf Tautenhahn

See Also

rawEIC,xcmsRaw-class

plotPeaks-methods *Plot a grid of a large number of peaks*

Description

Plot extracted ion chromatograms for many peaks simultaneously, indicating peak integration start and end points with vertical grey lines.

Arguments

object	the xcmsRaw object
peaks	matrix with peak information as produced by findPeaks
figs	two-element vector describing the number of rows and the number of columns of peaks to plot, if missing then an approximately square grid that will fit the number of peaks supplied
width	width of chromatogram retention time to plot for each peak

Details

This function is intended to help graphically analyze the results of peak picking. It can help estimate the number of false positives and improper integration start and end points. Its output is very compact and tries to waste as little space as possible. Each plot is labeled with rounded m/z and retention time separated by a space.

Methods

```
object = "xcmsRaw" plotPeaks(object, peaks, figs, width = 200)
```

See Also

xcmsRaw-class, findPeaks, split.screen

plotQC

Plot m/z and RT deviations for QC purposes without external reference data

Description

Use "democracy" to determine the average m/z and RT deviations for a grouped xcmsSet, and dependency on sample or absolute m/z

Usage

```
plotQC(object, sampNames, sampColors, sampOrder, what)
```

Arguments

object	A grouped xcmsSet
sampNames	Override sample names (e.g. with simplified names)
sampColors	Provide a set of colors (default: monochrome ?)
sampOrder	Override the order of samples, e.g. to bring them in order of measurement to detect time drift
what	A vector of which QC plots to generate. "mzdevhist": histogram of mz devia- tions. Should be gaussian shaped. If it is multimodal, then some peaks seem to have a systematically higher m/z deviation "rtdevhist": histogram of RT devi- ations. Should be gaussian shaped. If it is multimodal, then some peaks seem to have a systematically higher RT deviation "mzdevmass": Shows whether m/z deviations are absolute m/z dependent, could indicate miscalibration "mzdev- time": Shows whether m/z deviations are RT dependent, could indicate instru- ment drift "mzdevsample": median mz deviation for each sample, indicates out- liers "rtdevsample": median RT deviation for each sample, indicates out-

Details

plotQC() is a warpper to create a set of diagnostic plots. For the m/z deviations, the median of all m/z withon one group are assumed.

Value

No return value

Author(s)

Michael Wenk, Michael Wenk <michael.wenk@student.uni-halle.de>

Examples

```
library(faahKO)
xsg <- group(faahko)

plotQC(xsg, what="mzdevhist")
plotQC(xsg, what="rtdevhist")
plotQC(xsg, what="mzdevmass")
plotQC(xsg, what="mzdevtime")
plotQC(xsg, what="mzdevsample")
plotQC(xsg, what="rtdevsample")</pre>
```

plotRaw-methods

Description

Produce a scatterplot showing raw data point location in retention time and m/z. This plot is more useful for centroided data than continuum data.

Arguments

object	the xcmsRaw object
mzrange	numeric vector of length ≥ 2 whose range will be used to select the masses to plot
rtrange	numeric vector of length $>= 2$ whose range will be used to select the retention times to plot
scanrange	numeric vector of length ≥ 2 whose range will be used to select scans to plot
log	logical, log transform intensity
title	main title of the plot

Value

A matrix with the points plotted.

Methods

object = "xcmsRaw" plotRaw(object, mzrange = numeric(), rtrange = numeric(),

See Also

xcmsRaw-class

plotrt-methods Plot retention time deviation profiles

Description

Use corrected retention times for each sample to calculate retention time deviation profiles and plot each on the same graph.

object	the xcmsSet object
col	vector of colors for plotting each sample
ty	vector of line and point types for plotting each sample
leg	logical plot legend with sample labels
densplit	logical, also plot peak overall peak density

Methods

```
object = "xcmsSet" plotrt(object, col = NULL, ty = NULL, leg = TRUE, densplit = FALSE)
```

See Also

xcmsSet-class, retcor

plotScan-methods *Plot a single mass scan*

Description

Plot a single mass scan using the impulse representation. Most useful for centroided data.

Arguments

object	the xcmsRaw object
scan	integer with number of scan to plot
mzrange	numeric vector of length >= 2 whose range will be used to select masses to plot
ident	logical, use mouse to interactively identify and label individual masses

Methods

```
object = "xcmsRaw" plotScan(object, scan, mzrange = numeric(), ident = FALSE)
```

See Also

xcmsRaw-class

plotSpec-methods *Plot mass spectra from the profile matrix*

Description

Uses the pre-generated profile mode matrix to plot mass spectra over a specified retention time range.

Arguments

object	the xcmsRaw object
ident	logical, use mouse to identify and label peaks
vline	numeric vector with locations of vertical lines
	arguments passed to profRange

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plotSurf-methods

Value

If ident == TRUE, an integer vector with the indecies of the points that were identified. Otherwise a two-column matrix with the plotted points.

Methods

```
object = "xcmsRaw" plotSpec(object, ident = FALSE, vline = numeric(0), ...)
```

See Also

xcmsRaw-class

plotSurf-methods *Plot profile matrix 3D surface using OpenGL*

Description

This method uses the rgl package to create interactive three dimensional representations of the profile matrix. It uses the terrain color scheme.

Arguments

object	the xcmsRaw object
log	logical, log transform intensity
aspect	numeric vector with a spect ratio of the $\mbox{m/z},$ retention time and intensity components of the plot
	arguments passed to profRange

Details

The rgl package is still in development and imposes some limitations on the output format. A bug in the axis label code means that the axis labels only go from 0 to the aspect ratio constant of that axis. Additionally the axes are not labeled with what they are.

It is important to only plot a small portion of the profile matrix. Large portions can quickly overwhelm your CPU and memory.

Methods

```
object = "xcmsRaw" plotSurf(object, log = FALSE, aspect = c(1, 1, .5), ...)
```

See Also

xcmsRaw-class

plotTIC-methods Pa

Description

Plot chromatogram of total ion count. Optionally allow identification of target peaks and view-ing/identification of individual spectra.

Arguments

object	the xcmsRaw object
ident	logical, use mouse to identify and label chromatographic peaks
msident	logical, use mouse to identify and label spectral peaks

Value

If ident == TRUE, an integer vector with the indecies of the points that were identified. Otherwise a two-column matrix with the plotted points.

Methods

```
object = "xcmsRaw" plotTIC(object, ident = FALSE, msident = FALSE)
```

See Also

xcmsRaw-class

profMedFilt-methods *Median filtering of the profile matrix*

Description

Apply a median filter of given size to a profile matrix.

Arguments

object	the xcmsRaw object
massrad	number of m/z grid points on either side to use for median calculation
scanrad	number of scan grid points on either side to use for median calculation

Methods

```
object = "xcmsRaw" profMedFilt(object, massrad = 0, scanrad = 0)
```

See Also

xcmsRaw-class, medianFilter

profMethod-methods Get and set method for generating profile data

Description

These methods get and set the method for generating profile (matrix) data from raw mass spectral data. It can currently be bin, binlin, binlinbase, or intlin.

Methods

object = "xcmsRaw" profMethod(object)

See Also

xcmsRaw-class, profMethod, profBin, plotSpec, plotChrom, findPeaks

profRange-methods	Specify a subset of profile mode data	
proritange methodo	specify a subset of profile mode data	

Description

Specify a subset of the profile mode matrix given a mass, time, or scan range. Allow flexible user entry for other functions.

Arguments

object	the xcmsRaw object
mzrange	single numeric mass or vector of masses
rtrange	single numeric time (in seconds) or vector of times
scanrange	single integer scan index or vector of indecies
	arguments to other functions

Details

This function handles selection of mass/time subsets of the profile matrix for other functions. It allows the user to specify such subsets in a variety of flexible ways with minimal typing.

Because R does partial argument matching, mzrange, scanrange, and rtrange can be specified in short form using m=, s=, and t=, respectively. If both a scanrange and rtrange are specified, then the rtrange specification takes precedence.

When specifying ranges, you may either enter a single number or a numeric vector. If a single number is entered, then the closest single scan or mass value is selected. If a vector is entered, then the range is set to the range() of the values entered. That allows specification of ranges using shortened, slightly non-standard syntax. For example, one could specify 400 to 500 seconds using any of the following: t=c(400, 500), t=c(500, 400), or t=400:500. Use of the sequence operator (:) can save several keystrokes when specifying ranges. However, while the sequence operator works well for specifying integer ranges, fractional ranges do not always work as well.

Value

A list with the folloing items:

mzrange	numeric vector with start and end mass
masslab	textual label of mass range
massidx	integer vector of mass indecies
scanrange	integer vector with stat ane end scans
scanlab	textual label of scan range
scanidx	integer vector of scan range
rtrange	numeric vector of start and end times
timelab	textual label of time range

Methods

object = "xcmsRaw" profRange(object, mzrange = numeric(), rtrange = numeric(), scanrang

See Also

xcmsRaw-class

profStep-methods Get and set m/z step for generating profile data

Description

These methods get and set the m/z step for generating profile (matrix) data from raw mass spectral data. Smaller steps yield more precision at the cost of greater memory usage.

Methods

object = "xcmsRaw" profStep(object)

See Also

xcmsRaw-class, profMethod

Examples

```
## Not run:
library(faahKO)
cdfpath <- system.file("cdf", package = "faahKO")
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)
xset <- xcmsRaw(cdffiles[1])
xset
plotSurf(xset, mass=c(200,500))
```

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profStep(xset)<-0.1 ## decrease the bin size to get better resolution
plotSurf(xset, mass=c(200, 500))
##works nicer on high resolution data.</pre>

End(Not run)

rawEIC-methods Get extracted ion chromatograms for specified m/z range

Description

Generate extracted ion chromatogram for m/z values of interest. The raw data is used in contrast to getEIC which uses data from the profile matrix.

Arguments

object	xcmsRaw object
mzrange	m/z range for EIC
rtrange	retention time range for EIC
scanrange	scan range for EIC

Value

A list of :

scan	scan number
intensity	added intensity values

Methods

```
object = "xcmsRaw" rawEIC(object, mzrange = numeric(), rtrange = numeric(), scanrange = numeric()
```

Author(s)

Ralf Tautenhahn

See Also

xcmsRaw-class

rawMat-methods

Description

Returns a matrix with columns for time, m/z, and intensity that represents the raw data from a chromatography mass spectrometry experiment.

Arguments

object	The container of the raw data
mzrange	Subset by m/z range
rtrange	Subset by retention time range
scanrange	Subset by scan index range
log	Whether to log transform the intensities

Value

A numeric matrix with three columns: time, mz and intensity.

Methods

```
object = "xcmsRaw" rawMat(object, mzrange = numeric(), rtrange = numeric(), scanrange = numeric()
```

Author(s)

Michael Lawrence

See Also

plotRaw for plotting the raw intensities

retcor-methods Correct retention time from different samples

Description

To correct differences between retention times between different samples, a number of of methods exist in XCMS. retcor is the generic method.

object	<pre>xcmsSet-class object</pre>
method	Method to use for retention time correction. See details.
	Optional arguments to be passed along

retcor.obiwarp

Details

Different algorithms can be used by specifying them with the method argument. For example to use the approach described by Smith et al (2006) one would use: retcor(object, method="loess"). This is also the default.

Further arguments given by ... are passed through to the function implementing the method.

A character vector of *nicknames* for the algorithms available is returned by getOption("BioC")\$xcms\$retcor.methods. If the nickname of a method is called "loess", the help page for that specific method can be accessed with ?retcor.loess.

Value

An xcmsSet object with corrected retution times.

Methods

```
object = "xcmsSet" retcor(object, ...)
```

See Also

retcor.loess retcor.obiwarp xcmsSet-class,

retcor.obiwarp Align retention times across samples with Obiwarp

Description

Calculate retention time deviations for each sample. It is based on the code at http://obi-warp. sourceforge.net/. However, this function is able to align multiple samples, by a center-star strategy.

For the original publication see

Chromatographic Alignment of ESI-LC-MS Proteomics Data Sets by Ordered Bijective Interpolated Warping John T. Prince and, Edward M. Marcotte Analytical Chemistry 2006 78 (17), 6140-6152

object	the xcmsSet object
plottype	if deviation plot retention time deviation
profStep	step size (in m/z) to use for profile generation from the raw data files
center	the index of the sample all others will be aligned to. If center==NULL, the sample with the most peaks is chosen as default.
col	vector of colors for plotting each sample
ty	vector of line and point types for plotting each sample

response	Responsiveness of warping. 0 will give a linear warp based on start and end points. 100 will use all bijective anchors
distFunc	DistFunc function: cor (Pearson's R) or cor_opt (default, calculate only 10% diagonal band of distance matrix, better runtime), cov (covariance), prd (product), euc (Euclidean distance)
gapInit	Penalty for Gap opening, see below
gapExtend	Penalty for Gap enlargement, see below
factorDiag	Local weighting applied to diagonal moves in alignment.
factorGap	Local weighting applied to gap moves in alignment.
localAlignment	Local rather than global alignment
initPenalty	Penalty for initiating alignment (for local alignment only) Default: 0
	Default gap penalties: (gapInit, gapExtend) [by distFunc type]: 'cor' = '0.3,2.4' 'cov' = '0,11.7' 'prd' = '0,7.8' 'euc' = '0.9,1.8'

Value

An xcmsSet object

Methods

object = "xcmsSet" retcor(object, method="obiwarp", plottype = c("none", "deviation"), prof-Step=1, center=NULL, col = NULL, ty = NULL, response=1, distFunc="cor_opt", gapInit=NULL, gapExtend=NULL, factorDiag=2, factorGap=1, localAlignment=0, initPenalty=0)

See Also

xcmsSet-class,

retcor.peakgroups-methods

Align retention times across samples

Description

These two methods use "well behaved" peak groups to calculate retention time deviations for every time point of each sample. Use smoothed deviations to align retention times.

object	the xcmsSet object
missing	number of missing samples to allow in retention time correction groups
extra	number of extra peaks to allow in retention time correction correction groups
smooth	either "loess" for non-linear alignment or "linear" for linear alignment
span	degree of smoothing for local polynomial regression fitting

retexp

family	if gaussian fitting is by least-squares with no outlier removal, and if symmetric a re-descending M estimator is used with Tukey's biweight function, allowing outlier removal
plottype	if deviation plot retention time deviation points and regression fit, and if mdevden also plot peak overall peak density and retention time correction peak density
col	vector of colors for plotting each sample
ty	vector of line and point types for plotting each sample

Value

An xcmsSet object

Methods

```
object = "xcmsSet" retcor(object, missing = 1, extra = 1, smooth = c("loess", "linear"), space
```

See Also

xcmsSet-class, loess retcor.obiwarp

retexp	
recenp	

Set retention time window to a specified width

Description

Expands (or contracts) the retention time window in each row of a matrix as defined by the retmin and retmax columns.

Usage

retexp(peakrange, width = 200)

Arguments

peakrange	maxtrix with columns retmin and retmax
width	new width for the window

Value

The altered matrix.

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

getEIC

sampnames-methods Get sample names

Description

Return sample names for an object

Value

A character vector with sample names.

Methods

object = "xcmsEIC" sampnames(object)
object = "xcmsSet" sampnames(object)

See Also

xcmsSet-class, xcmsEIC-class

specDist-methods Distance methods for xcmsSet, xcmsRaw and xsAnnotate

Description

There are several methods for calculating a distance between two sets of peaks in xcms. specDist is the generic method.

Arguments

object	a xcmsSet or xcmsRaw.
method	Method to use for distance calculation. See details.
	mzabs, mzppm and parameters for the distance function

Details

Different algorithms can be used by specifying them with the method argument. For example to use the "meanMZmatch" approach with xcmsSet one would use: specDist(object, peakIDs1, peakIDs2, method="meanMZmThis is also the default.

Further arguments given by ... are passed through to the function implementing the method.

A character vector of *nicknames* for the algorithms available is returned by getOption("BioC")\$xcms\$specDist.methods. If the nickname of a method is called "meanMZmatch", the help page for that specific method can be accessed with ?specDist.meanMZmatch.

specDist.cosine

Value

mzabs	maximum absolute deviation for two matching peaks
mzppm	relative deviations in ppm for two matching peaks
symmetric	use symmetric pairwise m/z-matches only, or each match

Methods

object = "xcmsSet" specDist(object, peakIDs1, peakIDs2,...)

```
object = "xsAnnotate" specDist(object, PSpec1, PSpec2,...)
```

Author(s)

Joachim Kutzera, <jkutzer@ipb-halle.de>

specDist.cosine *a Distance function based on matching peaks*

Description

This method calculates the distance of two sets of peaks using the cosine-distance.

Usage

specDist.cosine(peakTable1, peakTable2, mzabs=0.001, mzppm=10, mzExp=0.6, intExp=3, nPdiff=2, nPmin=8

Arguments

peakTable1	a Matrix containing at least m/z-values, row must be called "mz"
peakTable2	the matrix for the other mz-values
mzabs	maximum absolute deviation for two matching peaks
mzppm	relative deviations in ppm for two matching peaks
symmetric	use symmetric pairwise m/z-matches only, or each match
mzExp	the exponent used for mz
intExp	the exponent used for intensity
nPdiff	the maximum nrow-difference of the two peaktables
nPmin	the minimum absolute sum of peaks from both praktables

Details

The result is the cosine-distance of the product from weighted factors of mz and intensity from matching peaks in the two peaktables. The factors are calculated as wFact = $mz^mzExp * int^intExp$. if no distance is calculated (for example because no matching peaks were found) the return-value is NA.

Methods

```
peakTable1 = "matrix", peakTable2 = "matrix"
```

specDist.cosine(peakTable1, peakTable2, mzabs = 0.001,

Author(s)

Joachim Kutzera, <jkutzer@ipb-halle.de>

specDist.meanMZmatch a Distance function based on matching peaks

Description

This method calculates the distance of two sets of peaks.

Usage

```
specDist.meanMZmatch(peakTable1, peakTable2, matchdist=1, matchrate=1, mzabs=0.001, mzppm=10, symmetic
```

Arguments

peakTable1	a Matrix containing at least m/z-values, row must be called "mz"
peakTable2	the matrix for the other mz-values
mzabs	maximum absolute deviation for two matching peaks
mzppm	relative deviations in ppm for two matching peaks
symmetric	use symmetric pairwise m/z-matches only, or each match
matchdist	the weight for value one (see details)
matchrate	the weight for value two

Details

The result of the calculation is a weighted sum of two values. Value one is the mean absolute difference of the matching peaks, value two is the relation of matching peaks and non matching peaks. if no distance is calculated (for example because no matching peaks were found) the return-value is NA.

Methods

peakTable1 = "matrix", peakTable2 = "matrix"

specDist.meanMZmatch(peakTable1, peakTable2, matchd

Author(s)

Joachim Kutzera, <jkutzer@ipb-halle.de>

specDist.peakCount-methods

a Distance function based on matching peaks

Description

This method calculates the distance of two sets of peaks by just returning the number of matching peaks (m/z-values).

Usage

```
specDist.peakCount(peakTable1, peakTable2, mzabs=0.001, mzppm=10, symmetric=FALSE)
```

Arguments

peakTable1	a Matrix containing at least m/z-values, row must be called "mz"
peakTable2	the matrix for the other mz-values
mzabs	maximum absolute deviation for two matching peaks
mzppm	relative deviations in ppm for two matching peaks
symmetric	use symmetric pairwise m/z-matches only, or each match

Methods

```
peakTable1 = "matrix", peakTable2 = "matrix" specDist.peakCount(peakTable1, peakTable2, mzppm=10,sy
```

Author(s)

Joachim Kutzera, <jkutzer@ipb-halle.de>

specNoise

Calculate noise for a sparse continuum mass spectrum

Description

Given a sparse continuum mass spectrum, determine regions where no signal is present, substituting half of the minimum intensity for those regions. Calculate the noise level as the weighted mean of the regions with signal and the regions without signal. If there is only one raw peak, return zero.

Usage

```
specNoise(spec, gap = quantile(diff(spec[, "mz"]), 0.9))
```

specPeaks

Arguments

spec	matrix with named columns mz and intensity
gap	threshold above which to data points are considerd to be separated by a blank
	region and not bridged by an interpolating line

Details

The default gap value is determined from the 90th percentile of the pair-wise differences between adjacent mass values.

Value

A numeric noise level

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

getSpec, specPeaks

specPeaks

Identify peaks in a sparse continuum mode spectrum

Description

Given a spectrum, identify and list significant peaks as determined by several criteria.

Usage

```
specPeaks(spec, sn = 20, mzgap = 0.2)
```

Arguments

spec	matrix with named columns mz and intensity
sn	minimum signal to noise ratio
mzgap	minimal distance between adjacent peaks, with smaller peaks being excluded

Details

Peaks must meet two criteria to be considered peaks: 1) Their s/n ratio must exceed a certain threshold. 2) They must not be within a given distance of any greater intensity peaks.

split.xcmsRaw

Value

A matrix with columns:

mz	m/z at maximum peak intensity
intensity	maximum intensity of the peak
fwhm	full width at half max of the peak

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

getSpec, specNoise

split.xcmsRaw Divide an xcmsRaw object

Description

Divides the scans from a xcmsRaw object into a list of multiple objects. MS\$^n\$ data is discarded.

Arguments

x	xcmsRaw object
f	factor such that factor(f) defines the scans which go into the new xcmsRaw objects
drop	logical indicating if levels that do not occur should be dropped (if 'f' is a 'factor' or a list).
	further potential arguments passed to methods.

Value

A list of xcmsRaw objects.

Methods

xr = "xcmsRaw" split(x, f, drop = TRUE, ...)

Author(s)

Steffen Neumann, <sneumann(at)ipb-halle.de>

See Also

xcmsRaw-class

split.xcmsSet

Description

Divides the samples and peaks from a xcmsSet object into a list of multiple objects. Group data is discarded.

Arguments

xs	xcmsSet object
f	factor such that factor(f) defines the grouping
drop	logical indicating if levels that do not occur should be dropped (if 'f' is a 'factor' or a list).
•••	further potential arguments passed to methods.

Value

A list of xcmsSet objects.

Methods

xs = "xcmsSet" split(x, f, drop = TRUE, ...)

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

xcmsSet-class

SSgauss

Gaussian Model

Description

This selfStart model evalueates the Gaussian model and its gradient. It has an initial attribute that will evalueate the initial estimates of the parameters mu, sigma, and h.

Usage

SSgauss(x, mu, sigma, h)

stitch-methods

Arguments

x	a numeric vector of values at which to evaluate the model
mu	mean of the distribution function
sigma	standard deviation of the distribution fuction
h	height of the distribution function

Details

Initial values for mu and h are chosen from the maximal value of x. The initial value for sigma is determined from the area under x divided by h*sqrt(2*pi).

Value

A numeric vector of the same length as x. It is the value of the expression $h \exp(-(x-mu)^2/(2*sigma^2))$, which is a modified gaussian function where the maximum height is treated as a separate parameter not dependent on sigma. If arguments mu, sigma, and h are names of objects, the gradient matrix with respect to these names is attached as an attribute named gradient.

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

nls, selfStart

stitch-methods Correct gaps in data

Description

Fixes gaps in data due to calibration scans or lock mass. Automatically detects file type and calls the relevant method. The mzXML file keeps the data the same length in time but overwrites the lock mass scans. The netCDF version adds the scans back into the data thereby increasing the length of the data and correcting for the unseen gap.

object	An xcmsRaw-class object
lockMass	A dataframe of locations of the gaps
freq	The intervals of the lock mass scans
start	The starting lock mass scan location, default is 1

Details

makeacqNum takes locates the gap using the starting lock mass scan and it's intervals. This data frame is then used in stitch to correct for the gap caused by the lock mass. Correction works by using scans from either side of the gap to fill it in.

Value

stitch A corrected xcmsRaw-class object makeacqNum A numeric vector of scan locations corresponding to lock Mass scans

Methods

```
object = "xcmsRaw" stitch(object, lockMass=numeric())
object = "xcmsRaw" makeacqNum(object, freq=numeric(), start=1)
```

Author(s)

Paul Benton, <hpaul.benton08@imperial.ac.uk>

Examples

```
## Not run: library(xcms)
library(faahKO) ## These files do not have this problem to correct for but just for an example
cdfpath <- system.file("cdf", package = "faahKO")</pre>
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)</pre>
xr<-xcmsRaw(cdffiles[1])</pre>
xr
##Lets assume that the lockmass starts at 1 and is every 100 scans
lockMass<-xcms:::makeacqNum(xr, freq=100, start=1)</pre>
## these are equcal
lockmass<-AutoLockMass(xr)</pre>
ob<-stitch(xr, lockMass)</pre>
ob
#plot the old data before correction
foo<-rawEIC(xr, m=c(200,210), scan=c(80,140))</pre>
plot(foo$scan, foo$intensity, type="h")
#plot the new corrected data to see what changed
foo<-rawEIC(ob, m=c(200,210), scan=c(80,140))</pre>
plot(foo$scan, foo$intensity, type="h")
## End(Not run)
```

verify.mzQuantM Verify an mzQuantML file

Description

Export in XML data formats: verify the written data

Usage

verify.mzQuantML(filename, xsdfilename)

Arguments

filename	filename (may include full path) for the output file. Pipes or URLs are not allowed.
xsdfilename	Filename of the XSD to verify against (may include full path)

Details

The verify.mzQuantML() function will verify an PSI standard format mzQuantML document against the XSD schemda, see http://www.psidev.info/mzquantml

Value

None.

See Also

write.mzQuantML

write.cdf-methods Save an xcmsRaw object to file

Description

Write the raw data to a (simple) CDF file.

Arguments

object	the xcmsRaw object
filename	filename (may include full path) for the CDF file. Pipes or URLs are not allowed.

Details

Currently the only application known to read the resulting file is XCMS. Others, especially those which build on the AndiMS library, will refuse to load the output.

Value

None.

Methods

object = "xcmsRaw" write.cdf(object, filename)

See Also

xcmsRaw-class, xcmsRaw,

write.mzdata-methods Save an xcmsRaw object to a file

Description

Write the raw data to a (simple) mzData file.

Arguments

object	the xcmsRaw object
filename	filename (may include full path) for the mzData file. Pipes or URLs are not allowed.

Details

This function will export a given xcmsRaw object to an mzData file. The mzData file will contain a <spectrumList> containing the <spectrum> with mass and intensity values in 32 bit precision. Other formats are currently not supported. Any header information (e.g. additional <software> information or <cvParams>) will be lost. Currently, also any MSn information will not be stored.

Value

None.

Methods

object = "xcmsRaw" write.mzdata(object, filename)

See Also

xcmsRaw-class, xcmsRaw,

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write.mzQuantML-methods

Save an xcmsSet object to an PSI mzQuantML file

Description

Export in XML data formats: Write the processed data in an xcmsSet to mzQuantML.

Arguments

object	the xcmsRaw or xcmsSet object
filename	filename (may include full path) for the output file. Pipes or URLs are not allowed.

Details

The write.mzQuantML() function will write a (grouped) xcmsSet into the PSI standard format mzQuantML, see http://www.psidev.info/mzquantml

Value

None.

Methods

object = "xcmsSet" write.mzQuantML(object, filename)

See Also

xcmsSet-class, xcmsSet, verify.mzQuantML,

xcmsEIC-class Class xcmsEIC, a class for multi-sample extracted ion chromatograms

Description

This class is used to store and plot parallel extracted ion chromatograms from multiple sample files. It integrates with the xcmsSet class to display peak area integrated during peak identification or fill-in.

Objects from the Class

Objects can be created with the getEIC method of the xcmsSet class. Objects can also be created by calls of the form new("xcmsEIC", ...).

eic: list containing named entries for every sample. for each entry, a list of two column EIC matricies with retention time and intensity

mzrange: two column matrix containing starting and ending m/z for each EIC

rtrange: two column matrix containing starting and ending time for each EIC

rt: either "raw" or "corrected" to specify retention times contained in the object

groupnames: group names from xcmsSet object used to generate EICs

Methods

groupnames signature(object = "xcmsEIC"): get groupnames slot mzrange signature(object = "xcmsEIC"): get mzrange slot plot signature(x = "xcmsEIC"): plot the extracted ion chromatograms rtrange signature(object = "xcmsEIC"): get rtrange slot sampnames signature(object = "xcmsEIC"): get sample names

Note

No notes yet.

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

getEIC

xcmsFileSource-class Base class for loading raw data from a file

Description

Data sources which read data from a file should inherit from this class. The xcms package provides classes to read from netCDF, mzData, mzXML, and mzML files using xcmsFileSource.

This class should be considered virtual and will not work if passed to loadRaw-methods. The reason it is not explicitly virtual is that there does not appear to be a way for a class to be both virtual and have a data part (which lets functions treat objects as if they were character strings).

This class validates that a file exists at the path given.

Objects from the Class

xcmsFileSource objects should not be instantiated directly. Instead, create subclasses and instantiate those.

xcmsFragments

Slots

.Data: Object of class "character". File path of a file from which to read raw data as the object's data part

Extends

Class "character", from data part. Class "xcmsSource", directly.

Methods

xcmsSource signature(object = "character"): Create an xcmsFileSource object referencing the given file name.

Author(s)

Daniel Hackney <dan@haxney.org>

See Also

xcmsSource

xcmsFragments

Constructor for xcmsFragments objects which holds Tandem MS peaks

Description

EXPERIMANTAL FEATURE

xcmsFragments is an object similar to xcmsSet, which holds peaks picked (or collected) from one or several xcmsRaw objects.

There are still discussions going on about the exact API for MS\$^n\$ data, so this is likely to change in the future. The code is not yet pipeline-ified.

Usage

xcmsFragments(xs, ...)

Arguments

XS	A xcmsSet-class object which contains picked ms1-peaks from one or several experiments
	further arguments to the collect method

Details

After running collect(xFragments,xSet) The peaktable of the xcmsFragments includes the ms1Peaks from all experiments stored in a xcmsSet-object. Further it contains the relevant MSn-peaks from the xcmsRaw-objects, which were created temporarily with the paths in xcmsSet.

Value

An xcmsFragments object.

Author(s)

Joachim Kutzera, Steffen Neumann, <sneumann@ipb-halle.de>

See Also

xcmsFragments-class, collect

xcmsFragments-class Class xcmsFragments, a class for handling Tandem MS and MS\$^n\$ data

Description

This class is similar to xcmsSet because it stores peaks from a number of individual files. However, xcmsFragments keeps Tandem MS and e.g. Ion Trap or Orbitrap MS\$^n\$ peaks, including the parent ion relationships.

Objects from the Class

Objects can be created with the xcmsFragments constructor and filled with peaks using the collect method.

Slots

- peaks: matrix with colmns peakID (MS1 parent in corresponding xcmsSet), MSnParentPeakID (parent peak within this xcmsFragments), msLevel (e.g. 2 for Tandem MS), rt (retention time in case of LC data), mz (fragment mass-to-charge), intensity (peak intensity extracted from the original xcmsSet), sample (the index of the rawData-file).
- MS2spec: This is a list of matrixes. Each matrix in the list is a single collected spectra from collect. The column ID's are mz, intensity, and full width half maximum(fwhm). The fwhm column is only relevant if the spectra came from profile data.
- specinfo: This is a matrix with reference data for the spectra in MS2spec. The column id's are preMZ, AccMZ, rtmin, rtmax, ref, CollisionEnergy. The preMZ is precursor mass from the MS1 scan. This mass is given by the XML file. With some instruments this mass is only given as nominal mass, therefore a AccMZ is given which is a weighted average mass from the MS1 scan of the collected spectra. The retention time is given by rtmin and rtmax. The ref column is a pointer to the MS2spec matrix spectra. The collisionEnergy column is the collision Energy for the spectra.

xcmsPapply

Methods

- collect signature(object = "xcmsFragments"): gets a xcmsSet-object, collects ms1-peaks
 from it and the msn-peaks from the corresponding xcmsRaw-files.
- plotTree signature(object = "xcmsFragments"): prints a (text based) pseudo-tree of the peaktable to display the dependencies of the peaks among each other.
- show signature(object = "xcmsFragments"): print a human-readable description of this object to the console.

Note

No notes yet.

Author(s)

S. Neumann, J. Kutzera

References

A parallel effort in metabolite profiling data sharing: http://metlin.scripps.edu/

xcmsPapply

See Also

xcmsRaw

xcmsPapply

Description

An apply-like function which uses Rmpi to distribute the processing evenly across a cluster. Will use a non-MPI version if distributed processing is not available.

Usage

Arguments

arg_sets	a list, where each item will be given as an argument to papply_action	
papply_action	A function which takes one argument. It will be called on each element of arg_sets	
papply_commondata		
	A list containing the names and values of variables to be accessible to the pap- ply_action. 'attach' is used locally to import this list.	

show_errors	If set to TRUE, overrides Rmpi's default, and messages for errors which occur in R slaves are produced.
do_trace	If set to TRUE, causes the papply_action function to be traced. i.e. Each state- ment is output before it is executed by the slaves.
also_trace	If supplied an array of function names, as strings, tracing will also occur for the specified functions.

Details

Similar to apply and lapply, applies a function to all items of a list, and returns a list with the corresponding results.

Uses Rmpi to implement a pull idiom in order to distribute the processing evenly across a cluster. If Rmpi is not available, or there are no slaves, implements this as a non-parallel algorithm.

xcmsPapply is a modified version of the papply function from package papply 0.2 (Duane Currie). Parts of the slave function were wrapped in try() to make it failsafe and progress output was added.

Make sure Rmpi was installed properly by executing the example below. Rmpi was tested with

- OpenMPI: Unix, http://www.open-mpi.org/, don't forget to export MPI_ROOT before installing Rmpi e.g. export MPI_ROOT=/usr/lib/openmpi
- DeinoMPI: Windows, http://mpi.deino.net/, also see http://www.stats.uwo.ca/faculty/ yu/Rmpi/

Value

A list of return values from papply_action. Each value corresponds to the element of arg_sets used as a parameter to papply_action

Note

Does not support distributing recursive calls in parallel. If papply is used inside papply_action, it will call a non-parallel version

Author(s)

Duane Currie <duane.currie@acadiau.ca>, modified by Ralf Tautenhahn <rtautenh@ipb-halle.de>.

References

http://ace.acadiau.ca/math/ACMMaC/software/papply/

Examples

```
## Not run:
library(Rmpi)
library(xcms)
```

number_lists <- list(1:10,4:40,2:27)</pre>

mpi.spawn.Rslaves(nslaves=2)

xcmsPeaks-class

```
results <- xcmsPapply(number_lists,sum)
results
mpi.close.Rslaves()
## End(Not run)</pre>
```

xcmsPeaks-class A matrix of peaks

Description

A matrix of peak information. The actual columns depend on how it is generated (i.e. the findPeaks method).

Objects from the Class

Objects can be created by calls of the form new("xcmsPeaks", ...).

Slots

.Data: The matrix holding the peak information

Extends

Class "matrix", from data part. Class "array", by class "matrix", distance 2. Class "structure", by class "matrix", distance 3. Class "vector", by class "matrix", distance 4, with explicit coerce.

Methods

None yet. Some utilities for working with peak data would be nice.

Author(s)

Michael Lawrence

See Also

findPeaks for detecting peaks in an xcmsRaw.

xcmsRaw

Description

This function handles the task of reading a NetCDF/mzXML file containing LC/MS or GC/MS data into a new xcmsRaw object. It also transforms the data into profile (maxrix) mode for efficient plotting and data exploration.

Usage

```
xcmsRaw(filename, profstep = 1, profmethod = "bin", profparam =
list(), includeMSn=FALSE, mslevel=NULL, scanrange=NULL)
```

deepCopy(object)

Arguments

filename	path name of the NetCDF or mzXML file to read
profstep	step size (in m/z) to use for profile generation
profmethod	method to use for profile generation
profparam	extra parameters to use for profile generation
includeMSn	only for XML file formats: also read MS\$^n\$ (Tandem-MS of Ion-/Orbi- Trap spectra)
mslevel	move data from mslevel into normal MS1 slots, e.g. for peak picking and visualisation
scanrange	scan range to read
object	An xcmsRaw object

Details

The scanrange to import can be restricted, otherwise all MS1 data is read. If profstep is set to 0, no profile matrix is generated. Unless includeMSn=TRUE only first level MS data is read, not MS/MS, etc.

deepCopy(xraw) will create a copy of the xcmsRaw object with its own copy of mz and intensity data in xraw@env.

Value

A xcmsRaw object.

Author(s)

Colin A. Smith, <csmith@scripps.edu>

xcmsRaw

References

NetCDF file format: http://my.unidata.ucar.edu/content/software/netcdf/ http://www.astm.org/Standards/E2077.htm http://www.astm.org/Standards/E2078.htm

mzXML file format: http://sashimi.sourceforge.net/software_glossolalia.html

PSI-MS working group who developed mzData and mzML file formats: http://www.psidev. info/index.php?q=node/80

Parser used for XML file formats: http://tools.proteomecenter.org/wiki/index.php?title= Software:RAMP

See Also

xcmsRaw-class, profStep, profMethod xcmsFragments

Examples

```
## Not run:
library(xcms)
library(faahKO)
cdfpath <- system.file("cdf", package = "faahKO")
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)
xr<-xcmsRaw(cdffiles[1])
xr
##This gives some information about the file
names(attributes(xr))
## Lets have a look at the structure of the object
```

```
str(xr)
##same but with a preview of each slot in the object
##S0... lets have a look at how this works
head(xr@scanindex)
#[1] 0 429 860 1291 1718 2140
xr@env$mz[425:430]
#[1] 596.3 597.0 597.3 598.1 599.3 200.1
##We can see that the 429 index is the last mz of scan 1 therefore...
```

```
mz.scan1<-xr@env$mz[(1+xr@scanindex[1]):xr@scanindex[2]]
intensity.scan1<-xr@env$intensity[(1+xr@scanindex[1]):xr@scanindex[2]]
plot(mz.scan1, intensity.scan1, type="h", main=paste("Scan 1 of file", basename(cdffiles[1]), sep=""))
##the easier way :p
scan1<-getScan(xr, 1)
head(scan1)
plotScan(xr, 1)</pre>
```

End(Not run)

```
xcmsRaw-class
```

Description

This class handles processing and visualization of the raw data from a single LC/MS or GS/MS run. It includes methods for producing a standard suite of plots including individual spectra, multi-scan average spectra, TIC, and EIC. It will also produce a feature list of significant peaks using matched filtration.

Objects from the Class

Objects can be created with the xcmsRaw constructor which reads data from a NetCDF file into a new object.

Slots

acquisitionNum: acquisitionNum

env: environment with three variables: mz - concatenated m/z values for all scans, intensity - corresponding signal intensity for each m/z value, and profile - matrix represention of the intensity values with columns representing scans and rows representing equally spaced m/z values

filepath: Path to the raw data file

- gradient: matrix with first row, time, containing the time point for interpolation and successive columns representing solvent fractions at each point
- msnAcquisitionNum: for each scan a unique acquisition number as reported via "spectrum id" (mzData) or "<scan num=...>" and "<scanOrigin num=...>" (mzXML)

msnCollisionEnergy: "CollisionEnergy" (mzData) or "collisionEnergy" (mzXML)

msnLevel: for each scan the "msLevel" (both mzData and mzXML)

msnPrecursorCharge: "ChargeState" (mzData) and "precursorCharge" (mzXML)

msnPrecursorIntensity: "Intensity" (mzData) or "precursorIntensity" (mzXML)

msnPrecursorMz: "MassToChargeRatio" (mzData) or "precursorMz" (mzXML)

msnPrecursorScan: "spectrumRef" (both mzData and mzXML)

msnRt: Retention time of the scan

msnScanindex: msnScanindex

- mzrange: numeric vector of length 2 with minimum and maximum m/z values represented in the profile matrix
- polarity: polarity

profmethod: characer value with name of method used for generating the profile matrix

- profparam: profparam
- scanindex: integer vector with starting positions of each scan in the mz and intensity variables
 (note that index values are based off a 0 initial position instead of 1)

scantime: numeric vector with acquisition time (in seconds) for each scan

- tic: numeric vector with total ion count (intensity) for each scan
- mslevel: Numeric representing the MS level that is present in MS1 slot. This slot should be accessed through its getter method mslevel.
- scanrange: Numeric of length 2 specifying the scan range (or NULL for the full range). This slot should be accessed through its getter method scanrange.

Methods

- findPeaks signature(object = "xcmsRaw"): feature detection using matched filtration in the chromatographic time domain
- **getEIC** signature(object = "xcmsRaw"): get extracted ion chromatograms in specified m/z ranges. This will return the total ion chromatogram (TIC) if the m/z range corresponds to the full m/z range (i.e. sum of all signals per retention time across all m/z).

getPeaks signature(object = "xcmsRaw"): get data for peaks in specified m/z and time ranges

- getScan signature(object = "xcmsRaw"): get m/z and intensity values for a single mass scan
- getSpec signature(object = "xcmsRaw"): get average m/z and intensity values for multiple
 mass scans

image signature(x = "xcmsRaw"): get data for peaks in specified m/z and time ranges

levelplot Create an image of the raw (profile) data m/z against retention time, with the intensity color coded.

mslevel Getter method for the mslevel slot.

- plotChrom signature(object = "xcmsRaw"): plot a chromatogram from profile data
- plotRaw signature(object = "xcmsRaw"): plot locations of raw intensity data points
- plotScan signature(object = "xcmsRaw"): plot a mass spectrum of an individual scan from the raw data
- plotSpec signature(object = "xcmsRaw"): plot a mass spectrum from profile data
- plotSurf signature(object = "xcmsRaw"): experimental method for plotting 3D surface of
 profile data with rgl.
- plotTIC signature(object = "xcmsRaw"): plot total ion count chromatogram
- profinfo signature(object = "xcmsRaw"): returns a list containing the profile generation method
 and step (profile m/z step size) and eventual additional parameters to the profile function.
- profMedFilt signature(object = "xcmsRaw"): median filter profile data in time and m/z dimensions
- profMethod<- signature(object = "xcmsRaw"): change the method of generating the profile
 matrix</pre>
- profMethod signature(object = "xcmsRaw"): get the method of generating the profile matrix
- profMz signature(object = "xcmsRaw"): get vector of m/z values for each row of the profile
 matrix
- profRange signature(object = "xcmsRaw"): interpret flexible ways of specifying subsets of
 the profile matrix

```
profStep<- signature(object = "xcmsRaw"): change the m/z step used for generating the
    profile matrix
profStep signature(object = "xcmsRaw"): get the m/z step used for generating the profile
    matrix
revMz signature(object = "xcmsRaw"): reverse the order of the data points for each scan
scanrange Getter method for the scanrange slot.
sortMz signature(object = "xcmsRaw"): sort the data points by increasing m/z for each scan
stitch signature(object = "xcmsRaw"): Raw data correction for lock mass calibration gaps.</pre>
```

Note

No notes yet.

Author(s)

Colin A. Smith, <csmith@scripps.edu>, Johannes Rainer <johannes.rainer@eurac.edu>

References

A parallel effort in metabolite profiling data sharing: http://metlin.scripps.edu/

See Also

xcmsRaw

xcmsSet	Constructor for xcmsSet objects which finds peaks in NetCDF/mzXML
	files

Description

This function handles the construction of xcmsSet objects. It finds peaks in batch mode and presorts files from subdirectories into different classes suitable for grouping.

Usage

xcmsSet

Arguments

files	path names of the NetCDF/mzXML files to read	
snames	sample names. By default the file name without extension is used.	
sclass	sample classes.	
phenoData	data.frame or AnnotatedDataFrame defining the sample names and classes and other sample related properties. If not provided, the argument sclass or the subdirectories in which the samples are stored will be used to specify sample grouping.	
profmethod	method to use for profile generation.	
profparam	parameters to use for profile generation.	
polarity	filter raw data for positive/negative scans	
lockMassFreq	Performs correction for Waters LockMass function	
mslevel	perform peak picking on data of given mslevel	
nSlaves	number of slaves/cores to be used for parallel peak detection. MPI is used if installed, otherwise the snow package is employed for multicore support. If none of the two packages is available it uses the parallel package for parallel processing on multiple CPUs of the current machine.	
progressCallback		
	function to be called, when progressInfo changes (useful for GUIs)	
scanrange	scan range to read	
	further arguments to the findPeaks method of the xcmsRaw class	

Details

The default values of the files, snames, sclass, and phenoData arguments cause the function to recursively search for readable files. The filename without extention is used for the sample name. The subdirectory path is used for the sample class. If the files contain both positive and negative spectra, the polarity can be selected explicitly. The default (NULL) is to read all scans.

If phenoData is provided, it is stored to the phenoData slot of the returned xcmsSet class. If that data.frame contains a column named "class", its content will be returned by the sampclass method and thus be used for the group/class assignment of the individual files (e.g. for peak group-ing etc.). For more details see the help of the xcmsSet-class.

The step size (in m/z) to use for profile generation can be submitted either using the profparam argument (e.g. profparam=list(step=0.1)) or by submitting step=0.1.

The feature/peak detection algorithm can be specified with the method argument which defaults to the "matchFilter" method (findPeaks.matchedFilter). Possible values are returned by getOption("BioC")\$xcms\$findPeaks.methods.

The lock mass correction allows for the lock mass scan to be added back in with the last working scan. This correction gives better reproducibility between sample sets.

Value

A xcmsSet object.

Note

The arguments profmethod and profparam have no influence on the feature/peak detection. The step size parameter step for the profile generation in the findPeaks.matchedFilter peak detection algorithm can be passed using the

Author(s)

Colin A. Smith, <csmith@scripps.edu>

See Also

xcmsSet-class, findPeaks, profStep, profMethod, xcmsPapply

xcmsSet-class

Class xcmsSet, a class for preprocessing peak data

Description

This class transforms a set of peaks from multiple LC/MS or GC/MS samples into a matrix of preprocessed data. It groups the peaks and does nonlinear retention time correction without internal standards. It fills in missing peak values from raw data. Lastly, it generates extracted ion chromatograms for ions of interest.

Details

The phenoData slot (and phenoData parameter in the xcmsSet function) is intended to contain a data.frame describing all experimental factors, i.e. the samples along with their properties. If this data.frame contains a column named "class", this will be returned by the sampclass method and will thus be used by all methods to determine the sample grouping/class assignment (e.g. to define the colors in various plots or for the group method).

The sampclass<- method adds or replaces the "class" column in the phenoData slot. If a data.frame is submitted to this method, the interaction of its columns will be stored into the "class" column.

Also, similar to other classes in Bioconductor, the \$ method can be used to directly access all columns in the phenoData slot (e.g. use xset\$name on a xcmsSet object called "xset" to extract the values from a column named "name" in the phenoData slot).

Objects from the Class

Objects can be created with the xcmsSet constructor which gathers peaks from a set NetCDF files. Objects can also be created by calls of the form new("xcmsSet", ...).

xcmsSet-class

Slots

peaks: matrix containing peak data

filled: a vector with peak indices of peaks which have been added by a fillPeaks method,

groups: matrix containing statistics about peak groups

groupidx: list containing indices of peaks in each group

phenoData: a data frame containing the experimental design factors

- rt: list containing two lists, raw and corrected, each containing retention times for every scan of every sample
- filepaths: character vector with absolute path name of each NetCDF file
- profinfo: list containing the values method profile generation method, and step profile m/z step size and eventual additional parameters to the profile function.

dataCorrection logical vector filled if the waters Lock mass correction parameter is used.

polarity: a string ("positive" or "negative" or NULL) describing whether only positive or negative scans have been used reading the raw data.

progressInfo: progress informations for some xcms functions (for GUI)

progressCallback: function to be called, when progressInfo changes (for GUI)

- mslevel: Numeric representing the MS level on which the peak picking was performed (by default on MS1). This slot should be accessed through its getter method mslevel.
- scanrange: Numeric of length 2 specifying the scan range (or NULL for the full range). This slot should be accessed through its getter method scanrange.

Methods

c signature("xcmsSet"): combine objects together

filepaths<- signature(object = "xcmsSet"): set filepaths slot</pre>

filepaths signature(object = "xcmsSet"): get filepaths slot

diffreport signature(object = "xcmsSet"): create report of differentially regulated ions including EICs

fillPeaks signature(object = "xcmsSet"): fill in peak data for groups with missing peaks

getEIC signature(object = "xcmsSet"): get list of EICs for each sample in the set

getXcmsRaw signature(object = "xcmsSet", sampleidx=1,profmethod=profMethod(object), profstep=profSt
read the raw data for one or more files in the xcmsSet and return it. The default parameters
will apply all settings used in the original xcmsSet call to generate the xcmsSet object to be
applied also to the raw data. Parameter sampleidx allows to specify which raw file(s) should
be loaded.

groupidx <- signature(object = "xcmsSet"): set groupidx slot

groupidx signature(object = "xcmsSet"): get groupidx slot

groupnames signature(object = "xcmsSet"): get textual names for peak groups

groups<- signature(object = "xcmsSet"): set groups slot</pre>

groups signature(object = "xcmsSet"): get groups slot

- groupval signature(object = "xcmsSet"): get matrix of values from peak data with a row for each peak group
- group signature(object = "xcmsSet"): find groups of peaks across samples that share similar m/z and retention times
- mslevel Getter method for the mslevel slot.

peaks<- signature(object = "xcmsSet"): set peaks slot</pre>

- peaks signature(object = "xcmsSet"): get peaks slot
- plotrt signature(object = "xcmsSet"): plot retention time deviation profiles
- profinfo<- signature(object = "xcmsSet"): set profinfo slot</pre>
- profinfo signature(object = "xcmsSet"): get profinfo slot
- profMethod signature(object = "xcmsSet"): extract the method used to generate the profile
 matrix.
- profStep signature(object = "xcmsSet"): extract the profile step used for the generation of the profile matrix.
- retcor signature(object = "xcmsSet"): use initial grouping of peaks to do nonlinear loess
 retention time correction
- sampclass<- signature(object = "xcmsSet"): Replaces the column "class" in the phenoData
 slot. See details for more information.</pre>
- sampclass signature(object = "xcmsSet"): Returns the content of the column "class" from the phenoData slot or, if not present, the interaction of the experimental design factors (i.e. of the phenoData data.frame). See details for more information.
- phenoData<- signature(object = "xcmsSet"): set the phenoData slot</pre>
- phenoData signature(object = "xcmsSet"): get the phenoData slot
- progressCallback<- signature(object = "xcmsSet"): set the progressCallback slot</pre>
- progressCallback signature(object = "xcmsSet"): get the progressCallback slot
- scanrange Getter method for the scanrange slot.
- sampnames<- signature(object = "xcmsSet"): set rownames in the phenoData slot</pre>
- sampnames signature(object = "xcmsSet"): get rownames in the phenoData slot
- **split** signature("xcmsSet"): divide the xcmsSet into a list of xcmsSet objects depending on the provided factor. Note that only peak data will be preserved, i.e. eventual peak grouping information will be lost.
- object\$name, object\$name<-value Access and set name column in phenoData
- object[, i] Conducts subsetting of a xcmsSet instance. Only subsetting on columns, i.e. samples, is supported. Subsetting is performed on all slots, also on groups and groupidx. Parameter i can be an integer vector, a logical vector or a character vector of sample names (matching sampnames).

Note

No notes yet.

xcmsSource-class

Author(s)

Colin A. Smith, <csmith@scripps.edu>, Johannes Rainer <johannes.rainer@eurac.edu>

References

A parallel effort in metabolite profiling data sharing: http://metlin.scripps.edu/

See Also

xcmsSet

xcmsSource-class Virtual class for raw data sources

Description

This virtual class provides an implementation-independent way to load mass spectrometer data from various sources for use in an xcmsRaw object. Subclasses can be defined to enable data to be loaded from user-specified sources. The virtual class xcmsFileSource is included out of the box which contains a file name as a character string.

When implementing child classes of xcmsSource, a corresponding loadRaw-methods method must be provided which accepts the xcmsSource child class and returns a list in the format described in loadRaw-methods.

Objects from the Class

A virtual Class: No objects may be created from it.

Author(s)

Daniel Hackney, <dan@haxney.org>

See Also

xcmsSource-methods for creating xcmsSource objects in various ways.

xcmsSource-methods Create an xcmsSource object in a flexible way

Description

Users can define alternate means of reading data for xcmsRaw objects by creating new implementations of this method.

Methods

signature(object = "xcmsSource") Pass the object through unmodified.

Author(s)

Daniel Hackney, <dan@haxney.org>

See Also

xcmsSource

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