Package 'xcms'

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Title LC-MS and GC-MS Data Analysis

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.

Depends R (>= 4.0.0), BiocParallel (>= 1.8.0), MSnbase (>= 2.19.1)

- Imports mzR (>= 2.25.3), methods, Biobase, BiocGenerics, ProtGenerics (>= 1.25.1), lattice, RColorBrewer, plyr, RANN, MassSpecWavelet (>= 1.5.2), S4Vectors, robustbase, IRanges, SummarizedExperiment, MsCoreUtils, MsFeatures
- Suggests BiocStyle, caTools, knitr (>= 1.1.0), faahKO, msdata (>= 0.25.1), ncdf4, testthat, pander, magrittr, rmarkdown, multtest, MALDIquant, pheatmap, Spectra (>= 1.1.17), MsBackendMgf, progress
- Enhances Rgraphviz, rgl, XML

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URL https://github.com/sneumann/xcms

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R topics documented:

absent-methods
adjustRtime
adjustRtime-obiwarp
adjustRtime-peakGroups 13
applyAdjustedRtime
AutoLockMass-methods
bin,XCMSnExp-method 21
binYonX
breaks_on_binSize
breaks_on_nBins
c-methods
CalibrantMassParam-class
$calibrate-methods \ . \ . \ . \ . \ . \ . \ . \ . \ . \ $
$chromatogram, XCMSnExp-method \dots \dots \dots \dots \dots \dots \dots 32$

chromatographic-peak-detection	
chromPeakSpectra	. 36
CleanPeaksParam	. 38
collect-methods	. 40
correlate, Chromatogram, Chromatogram-method	. 41
diffreport-methods	
dirname	
do_adjustRtime_peakGroups	
do_findChromPeaks_centWave	
do_findChromPeaks_centWaveWithPredIsoROIs	
do_findChromPeaks_massifquant	
do_findChromPeaks_matchedFilter	
do_findPeaks_MSW	
do_groupChromPeaks_density	
do_groupChromPeaks_nearest	
do_groupPeaks_mzClust	
estimatePrecursorIntensity	
•	
etg	
exportMetaboAnalyst	
extractMsData,OnDiskMSnExp-method	
feature-grouping	
featureChromatograms	
featureSpectra	
featureSummary	
FillChromPeaksParam-class	
fillPeaks-methods	
fillPeaks.chrom-methods	
fillPeaks.MSW-methods	
filterColumnsIntensityAbove,MChromatograms-method	
filterFeatureDefinitions	. 90
FilterIntensityParam	. 94
findChromPeaks,Chromatogram,CentWaveParam-method	. 96
findChromPeaks,Chromatogram,MatchedFilterParam-method	. 98
findChromPeaks-centWave	. 99
findChromPeaks-centWaveWithPredIsoROIs	. 105
findChromPeaks-massifquant	. 109
findChromPeaks-matchedFilter	
findChromPeaksIsolationWindow	
findMZ	
findneutral	
findPeaks-methods	
findPeaks-MSW	
findPeaks.addPredictedIsotopeFeatures-methods	
findPeaks.centWave-methods	
findPeaks.centWaveWithPredictedIsotopeROIs-methods	
findPeaks.massifquant-methods	
findPeaks.matchedFilter,xcmsRaw-method	
findPeaks.MS1-methods	. 142

findPeaks.MSW,xcmsRaw-method	. 144
GenericParam-class	. 145
getEIC-methods	. 146
getPeaks-methods	. 147
getScan-methods	. 148
getSpec-methods	. 148
getXcmsRaw-methods	. 149
group-methods	. 150
group.density	. 151
group.mzClust	
group.nearest	
groupChromPeaks	
groupChromPeaks-density	
groupChromPeaks-mzClust	
groupChromPeaks-nearest	
groupFeatures-abundance-correlation	
groupFeatures-eic-similarity	
groupFeatures-similar-rtime	
groupnames,XCMSnExp-method	
groupnames-methods	
groupOverlaps	
groupval-methods	
highlightChromPeaks	
image-methods	
imputeLinInterpol	
imputeRowMin	
imputeRowMinRand	
isolationWindowTargetMz,OnDiskMSnExp-method	
levelplot-methods	
loadRaw-methods	
manualChromPeaks	
medianFilter	
MergeNeighboringPeaksParam	
msn2xcmsRaw	
overlappingFeatures	
peakPlots-methods	
peaksWithCentWave	
peaksWithMatchedFilter	
peakTable-methods	
phenoDataFromPaths	
plot.xcmsEIC	
plotAdjustedRtime	
1 5	
plotChrom-methods	
plotChromatogramsOverlay	
plotChromPeakDensity,XCMSnExp-method	
plotChromPeaks	
plotEIC-methods	
plotFeatureGroups	. 209

plotMsData	. 210
plotPeaks-methods	. 211
plotQC	. 212
plotRaw-methods	. 213
plotrt-methods	. 214
plotScan-methods	. 215
plotSpec-methods	. 215
plotSurf-methods	. 216
plotTIC-methods	. 217
ProcessHistory-class	. 217
profMat-xcmsSet	. 219
profMedFilt-methods	. 221
profMethod-methods	. 221
profRange-methods	. 222
profStep-methods	. 223
quantify,XCMSnExp-method	. 224
rawEIC-methods	
rawMat-methods	. 227
reconstructChromPeakSpectra	
removeIntensity,Chromatogram-method	
retcor-methods	
retcor.obiwarp	
retcor.peakgroups-methods	
retexp	
rla	
sampnames-methods	
showError,xcmsSet-method	
specDist-methods	
specDist.cosine	
specDist.meanMZmatch	
specDist.peakCount-methods	
specNoise	
specPeaks	
split.xcmsRaw	
split.xcmsSet	
SSgauss	. 243
stitch-methods	
updateObject,xcmsSet-method	
useOriginalCode	
verify.mzQuantM	
write.cdf-methods	
write.mzdata-methods	
write.mzQuantML-methods	
writeMSData,XCMSnExp,character-method	
writeMzTab	
XChromatograms	
xcms-deprecated	
xcmsEIC-class	

absent-methods

xcmsFileSource-class	266
xcmsFragments	267
xcmsFragments-class	268
XCMSnExp-class	269
xcmsPeaks-class	280
xcmsRaw	281
xcmsRaw-class	283
xcmsSet	286
xcmsSet-class	288
xcmsSource-class	290
xcmsSource-methods	291
xdata	291
[,xcmsRaw,logicalOrNumeric,missing,missing-method	292
	294

Index

absent-methods

Determine which peaks are absent / present in a sample class

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

adjustRtime

Description

The adjustRtime method(s) perform retention time correction (alignment) between chromatograms of different samples. These methods are part of the modernized xcms user interface.

The implemented retention time adjustment methods are:

- **peakGroups** retention time correction based on alignment of features (peak groups) present in most/all samples. See adjustRtime-peakGroups for more details.
- **obiwarp** alignment based on the complete mz-rt data. This method does not require any identified peaks or defined features. See adjustRtime-obiwarp for more details.

Author(s)

Johannes Rainer

See Also

retcor for the *old* retention time correction methods. plotAdjustedRtime for visualization of alignment results.

Other retention time correction methods: adjustRtime-obiwarp, adjustRtime-peakGroups

adjustRtime-obiwarp Align retention times across samples using Obiwarp

Description

This method performs retention time adjustment using the Obiwarp method [Prince 2006]. It is based on the code at http://obi-warp.sourceforge.net but supports alignment of multiple samples by aligning each against a *center* sample. The alignment is performed directly on the profile-matrix and can hence be performed independently of the peak detection or peak grouping.

It is also possible to exclude certain samples within an experiment from the estimation of the alignment models. The parameter subset allows to define the indices of samples within object that should be aligned. Samples not part of this subset are left out in the estimation of the alignment models, but their retention times are subsequently adjusted based on the alignment results of the closest sample in subset (close in terms of position within the object). Alignment could thus be performed on only *real* samples leaving out e.g. blanks, which are then in turn adjusted based on the closest real sample. Here it is up to the user to ensure that the samples within object are ordered correctly (e.g. by injection index).

How the non-subset samples are adjusted bases also on the parameter subsetAdjust: with subsetAdjust = "previous", each non-subset sample is adjusted based on the closest previous subset sample

which results in most cases with adjusted retention times of the non-subset sample being identical to the subset sample on which the adjustment bases. The second, default, option is to use subsetAdjust = "average" in which case each non subset sample is adjusted based on the average retention time adjustment from the previous and following subset sample. For the average a weighted mean is used with weights being the inverse of the distance of the non-subset sample to the subset samples used for alignment.

See also section Alignment of experiments including blanks in the xcms vignette for an example.

The ObiwarpParam class allows to specify all settings for the retention time adjustment based on the *obiwarp* method. Class Instances should be created using the ObiwarpParam constructor.

binSize,binSize<-: getter and setter for the binSize slot of the object.

centerSample,centerSample<-: getter and setter for the centerSample slot of the object.

response, response <-: getter and setter for the response slot of the object.

distFun,distFun<-: getter and setter for the distFun slot of the object.

gapInit,gapInit<-: getter and setter for the gapInit slot of the object.

gapExtend,gapExtend<-: getter and setter for the gapExtend slot of the object.

factorDiag, factorDiag<-: getter and setter for the factorDiag slot of the object.

factorGap,factorGap<-: getter and setter for the factorGap slot of the object.

localAlignment,localAlignment<-: getter and setter for the localAlignment slot of the object.

initPenalty, initPenalty <-: getter and setter for the initPenalty slot of the object.

subset, subset <-: getter and setter for the subset slot of the object.

subsetAdjust,subsetAdjust<-: getter and setter for the subsetAdjust slot of the object.

adjustRtime,XCMSnExp,ObiwarpParam: performs retention time correction/alignment based on the total mz-rt data using the *obiwarp* method.

Usage

```
ObiwarpParam(
  binSize = 1,
  centerSample = integer(),
  response = 1L,
  distFun = "cor_opt",
  gapInit = numeric(),
  gapExtend = numeric(),
  factorDiag = 2,
  factorGap = 1,
  localAlignment = FALSE,
  initPenalty = 0,
  subset = integer(),
  subsetAdjust = c("average", "previous")
)
## S4 method for signature 'OnDiskMSnExp,ObiwarpParam'
```

8

adjustRtime-obiwarp

```
## S4 method for signature 'ObiwarpParam'
binSize(object)
## S4 replacement method for signature 'ObiwarpParam'
binSize(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
centerSample(object)
## S4 replacement method for signature 'ObiwarpParam'
centerSample(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
response(object)
## S4 replacement method for signature 'ObiwarpParam'
response(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
distFun(object)
## S4 replacement method for signature 'ObiwarpParam'
distFun(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
gapInit(object)
## S4 replacement method for signature 'ObiwarpParam'
gapInit(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
gapExtend(object)
## S4 replacement method for signature 'ObiwarpParam'
gapExtend(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
factorDiag(object)
## S4 replacement method for signature 'ObiwarpParam'
factorDiag(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
factorGap(object)
## S4 replacement method for signature 'ObiwarpParam'
factorGap(object) <- value</pre>
```

```
## S4 method for signature 'ObiwarpParam'
localAlignment(object)
## S4 replacement method for signature 'ObiwarpParam'
localAlignment(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
initPenalty(object)
## S4 replacement method for signature 'ObiwarpParam'
initPenalty(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
subset(x)
## S4 replacement method for signature 'ObiwarpParam'
subset(object) <- value</pre>
## S4 method for signature 'ObiwarpParam'
subsetAdjust(object)
## S4 replacement method for signature 'ObiwarpParam'
subsetAdjust(object) <- value</pre>
## S4 method for signature 'XCMSnExp,ObiwarpParam'
adjustRtime(object, param, msLevel = 1L)
```

Arguments

binSize	numeric(1) defining the bin size (in mz dimension) to be used for the <i>profile matrix</i> generation. See step parameter in profile-matrix documentation for more details.
centerSample	integer(1) defining the index of the center sample in the experiment. It defaults to floor(median(1:length(fileNames(object)))). Note that if subset is used, the index passed with centerSample is within these subset samples.
response	numeric(1) defining the <i>responsiveness</i> of warping with response = 0 giving linear warping on start and end points and response = 100 warping using all bijective anchors.
distFun	character defining the distance function to be used. Allowed values are "cor" (Pearson's correlation), "cor_opt" (calculate only 10% diagonal band of distance matrix; better runtime), "cov" (covariance), "prd" (product) and "euc" (Euclidian distance). The default value is distFun = "cor_opt".
gapInit	<pre>numeric(1) defining the penalty for gap opening. The default value for gapInit depends on the value of distFun: for distFun = "cor" and distFun = "cor_opt" it is 0.3, for distFun = "cov" and distFun = "prd" 0.0 and for distFun = "euc" 0.9.</pre>
gapExtend	numeric(1) defining the penalty for gap enlargement. The default value for gapExtend depends on the value of distFun, for distFun = "cor" and distFun

10

	= "cor_opt" it is 2.4, for distFun = "cov" 11.7, for distFun = "euc" 1.8 and for distFun = "prd" 7.8.
factorDiag	numeric(1) defining the local weight applied to diagonal moves in the alignment.
factorGap	numeric(1) defining the local weight for gap moves in the alignment.
localAlignment	logical(1) whether a local alignment should be performed instead of the de- fault global alignment.
initPenalty	numeric(1) defining the penalty for initiating an alignment (for local alignment only).
subset	integer with the indices of samples within the experiment on which the align- ment models should be estimated. Samples not part of the subset are adjusted based on the closest subset sample. See description above for more details.
subsetAdjust	character specifying the method with which non-subset samples should be adjusted. Supported options are "previous" and "average" (default). See description above for more information.
object	For adjustRtime: an XCMSnExp object. For all other methods: a ObiwarpParam object.
param	A ObiwarpParam object containing all settings for the alignment method.
msLevel	integer defining the MS level on which the retention time should be performed.
value	The value for the slot.
х	a PeakGroupsParam object.

Value

The ObiwarpParam function returns a ObiwarpParam class instance with all of the settings specified for obiwarp retention time adjustment and alignment.

For adjustRtime,XCMSnExp,ObiwarpParam: a XCMSnExp object with the results of the retention time adjustment step. These can be accessed with the adjustedRtime method. Retention time correction does also adjust the retention time of the identified chromatographic peaks (accessed *via* chromPeaks. Note that retention time correction drops all previous peak grouping results from the result object.

For adjustRtime, OnDiskMSnExp, ObiwarpParam: a numeric with the adjusted retention times per spectra (in the same order than rtime).

Slots

binSize,centerSample,response,distFun,gapInit,gapExtend,factorDiag,factorGap,localAlignment,initPen See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the retcor methods. All of the settings to the alignment algorithm can be passed with a ObiwarpParam object.

Alignment using obiwarp is performed on the retention time of spectra of on MS level. Retention times for spectra of other MS levels are subsequently adjusted based on the adjustment function defined on the retention times of the spectra of MS level msLevel.

Calling adjustRtime on an XCMSnExp object will cause all peak grouping (correspondence) results and any previous retention time adjustment results to be dropped.

Author(s)

Colin Smith, Johannes Rainer

References

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

John T. Prince and Edward M. Marcotte. "Chromatographic Alignment of ESI-LC-MS Proteomic Data Sets by Ordered Bijective Interpolated Warping" *Anal. Chem.* 2006, 78 (17), 6140-6152.

See Also

retcor.obiwarp for the old user interface. plotAdjustedRtime for visualization of alignment results.

XCMSnExp for the object containing the results of the alignment.

Other retention time correction methods: adjustRtime-peakGroups, adjustRtime()

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Perform retention time correction:
res <- adjustRtime(faahko_sub, param = ObiwarpParam())
## As a result we get a numeric vector with the adjusted retention times for
## all spectra.
head(res)
## We can split this by file to get the adjusted retention times for each
## file
resL <- split(res, fromFile(res))</pre>
```

adjustRtime-peakGroups

Retention time correction based on alignment of house keeping peak groups

Description

This method performs retention time adjustment based on the alignment of chromatographic peak groups present in all/most samples (hence corresponding to house keeping compounds). First the retention time deviation of these peak groups is described by fitting either a polynomial (smooth = "loess") or a linear (smooth = "linear") model to the data points. These models are subsequently used to adjust the retention time of each spectrum in each sample.

It is also possible to exclude certain samples within an experiment from the estimation of the alignment models. The parameter subset allows to define the indices of samples within object that should be aligned. Samples not part of this subset are left out in the estimation of the alignment models, but their retention times are subsequently adjusted based on the alignment results of the closest sample in subset (close in terms of position within the object). Alignment could thus be performed on only *real* samples leaving out e.g. blanks, which are then in turn adjusted based on the closest real sample. Here it is up to the user to ensure that the samples within object are ordered correctly (e.g. by injection index).

How the non-subset samples are adjusted bases also on the parameter subsetAdjust: with subsetAdjust = "previous", each non-subset sample is adjusted based on the closest previous subset sample which results in most cases with adjusted retention times of the non-subset sample being identical to the subset sample on which the adjustment bases. The second, default, option is to use subsetAdjust = "average" in which case each non subset sample is adjusted based on the average retention time adjustment from the previous and following subset sample. For the average a weighted mean is used with weights being the inverse of the distance of the non-subset sample to the subset samples used for alignment.

See also section Alignment of experiments including blanks in the xcms vignette for an example.

The PeakGroupsParam class allows to specify all settings for the retention time adjustment based on *house keeping* peak groups present in most samples. Instances should be created with the PeakGroupsParam constructor.

adjustRtimePeakGroups returns the features (peak groups) which would, depending on the provided PeakGroupsParam, be selected for alignment/retention time correction.

minFraction,minFraction<-: getter and setter for the minFraction slot of the object.

extraPeaks, extraPeaks<-: getter and setter for the extraPeaks slot of the object.

smooth, smooth <-: getter and setter for the smooth slot of the object.

span, span<-: getter and setter for the span slot of the object.

family, family<-: getter and setter for the family slot of the object.

peakGroupsMatrix,peakGroupsMatrix<-: getter and setter for the peakGroupsMatrix slot of the object.

subset, subset <-: getter and setter for the subset slot of the object.

subsetAdjust,subsetAdjust<-: getter and setter for the subsetAdjust slot of the object.

adjustRtime,XCMSnExp,PeakGroupsParam: performs retention time correction based on the alignment of peak groups (features) found in all/most samples. The correction function identified on these peak groups is applied to the retention time of all spectra in the object, i.e. retention times of all spectra, also MS level > 1 are adjusted.

Usage

```
PeakGroupsParam(
  minFraction = 0.9,
  extraPeaks = 1,
  smooth = "loess",
  span = 0.2,
  family = "gaussian",
  peakGroupsMatrix = matrix(nrow = 0, ncol = 0),
  subset = integer(),
  subsetAdjust = c("average", "previous")
)
adjustRtimePeakGroups(object, param = PeakGroupsParam(), msLevel = 1L)
## S4 method for signature 'PeakGroupsParam'
minFraction(object)
## S4 replacement method for signature 'PeakGroupsParam'
minFraction(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
extraPeaks(object)
## S4 replacement method for signature 'PeakGroupsParam'
extraPeaks(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
smooth(x)
## S4 replacement method for signature 'PeakGroupsParam'
smooth(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
span(object)
## S4 replacement method for signature 'PeakGroupsParam'
span(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
family(object)
```

14

```
## S4 replacement method for signature 'PeakGroupsParam'
family(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
peakGroupsMatrix(object)
## S4 replacement method for signature 'PeakGroupsParam'
peakGroupsMatrix(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
subset(x)
## S4 replacement method for signature 'PeakGroupsParam'
subset(object) <- value</pre>
## S4 method for signature 'PeakGroupsParam'
subsetAdjust(object)
## S4 replacement method for signature 'PeakGroupsParam'
subsetAdjust(object) <- value</pre>
## S4 method for signature 'XCMSnExp,PeakGroupsParam'
adjustRtime(object, param, msLevel = 1L)
```

Arguments

minFraction	numeric(1) between 0 and 1 defining the minimum required fraction of samples in which peaks for the peak group were identified. Peak groups passing this criteria will aligned across samples and retention times of individual spectra will be adjusted based on this alignment. For minFraction = 1 the peak group has to contain peaks in all samples of the experiment. Note that if subset is provided, the specified fraction is relative to the defined subset of samples and not to the total number of samples within the experiment (i.e. a peak has to be present in the specified proportion of subset samples).
extraPeaks	numeric(1) defining the maximal number of additional peaks for all samples to be assigned to a peak group (i.e. feature) for retention time correction. For a data set with 6 samples, extraPeaks = 1 uses all peak groups with a total peak count <= 6 + 1. The total peak count is the total number of peaks being assigned to a peak group and considers also multiple peaks within a sample being assigned to the group.
smooth	character defining the function to be used, to interpolate corrected retention times for all peak groups. Either "loess" or "linear".
span	<pre>numeric(1) defining the degree of smoothing (if smooth = "loess"). This pa- rameter is passed to the internal call to loess.</pre>
family	character defining the method to be used for loess smoothing. Allowed values are "gaussian" and "symmetric".See loess for more information.

peakGroupsMatrix		
	optional matrix of (raw) retention times for the peak groups on which the align- ment should be performed. Each column represents a sample, each row a fea- ture/peak group. Such a matrix is for example returned by the adjustRtimePeakGroups method.	
subset	integer with the indices of samples within the experiment on which the align- ment models should be estimated. Samples not part of the subset are adjusted based on the closest subset sample. See description above for more details.	
subsetAdjust	character specifying the method with which non-subset samples should be adjusted. Supported options are "previous" and "average" (default). See description above for more information.	
object	For adjustRtime: an XCMSnExp object containing the results from a previous chromatographic peak detection (see findChromPeaks) and alignment analysis (see groupChromPeaks).	
	For all other methods: a PeakGroupsParam object.	
param	A PeakGroupsParam object containing all settings for the retention time correc- tion method	
msLevel	integer(1) specifying the MS level. Currently only MS level 1 is supported.	
value	The value for the slot.	
х	a PeakGroupsParam object.	

Value

The PeakGroupsParam function returns a PeakGroupsParam class instance with all of the settings specified for retention time adjustment based on *house keeping* features/peak groups.

For adjustRtimePeakGroups: a matrix, rows being features, columns samples, of retention times. The features are ordered by the median retention time across columns.

For adjustRtime: a XCMSnExp object with the results of the retention time adjustment step. These can be accessed with the adjustedRtime method. Retention time correction does also adjust the retention time of the identified chromatographic peaks (accessed *via* chromPeaks. Note that retention time correction drops all previous alignment results from the result object.

Slots

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the group methods. All of the settings to the alignment algorithm can be passed with a PeakGroupsParam object.

The matrix with the (raw) retention times of the peak groups used in the alignment is added to the peakGroupsMatrix slot of the PeakGroupsParam object that is stored into the corresponding *process history step* (see processHistory for how to access the process history).

adjustRtime-peakGroups

adjustRtimePeakGroups is supposed to be called *before* the sample alignment, but after a correspondence (peak grouping).

This method requires that a correspondence analysis has been performed on the data, i.e. that grouped chromatographic peaks/features are present (see groupChromPeaks for details).

Calling adjustRtime on an XCMSnExp object will cause all peak grouping (correspondence) results and any previous retention time adjustments to be dropped. In some instances, the adjustRtime, XCMSnExp, PeakGroupsPara re-adjusts adjusted retention times to ensure them being in the same order than the raw (original) retention times.

Author(s)

Colin Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" *Anal. Chem.* 2006, 78:779-787.

See Also

The do_adjustRtime_peakGroups core API function and retcor.peakgroups for the old user interface. plotAdjustedRtime for visualization of alignment results.

XCMSnExp for the object containing the results of the alignment.

Other retention time correction methods: adjustRtime-obiwarp, adjustRtime()

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")</pre>
res <- faahko_sub
## Disable parallel processing for this example
register(SerialParam())
head(chromPeaks(res))
## The number of peaks identified per sample:
table(chromPeaks(res)[, "sample"])
## Performing the peak grouping using the "peak density" method.
p \le PeakDensityParam(sampleGroups = c(1, 1, 1))
res <- groupChromPeaks(res, param = p)</pre>
## Perform the retention time adjustment using peak groups found in both
## files.
fgp <- PeakGroupsParam(minFraction = 1)</pre>
```

Before running the alignment we can evaluate which features (peak groups)

```
## would be used based on the specified parameters.
pkGrps <- adjustRtimePeakGroups(res, param = fgp)</pre>
## We can also plot these to evaluate if the peak groups span a large portion
## of the retention time range.
plot(x = pkGrps[, 1], y = rep(1, nrow(pkGrps)), xlim = range(rtime(res)),
   ylim = c(1, 2), xlab = "rt", ylab = "", yaxt = "n")
points(x = pkGrps[, 2], y = rep(2, nrow(pkGrps)))
segments(x0 = pkGrps[, 1], x1 = pkGrps[, 2],
   y0 = rep(1, nrow(pkGrps)), y1 = rep(2, nrow(pkGrps)))
grid()
axis(side = 2, at = c(1, 2, 3), labels = colnames(pkGrps))
## Next we perform the alignment.
res <- adjustRtime(res, param = fgp)</pre>
## Any grouping information was dropped
hasFeatures(res)
## Plot the raw against the adjusted retention times.
plot(rtime(res, adjusted = FALSE),
    rtime(res), pch = 16, cex = 0.25, col = fromFile(res))
## Adjusterd retention times can be accessed using
## rtime(object, adjusted = TRUE) and adjustedRtime
all.equal(rtime(res), adjustedRtime(res))
## To extract the retention times grouped by sample/file:
rts <- rtime(res, bySample = TRUE)</pre>
```

applyAdjustedRtime Replace raw with adjusted retention times

Description

Replaces the raw retention times with the adjusted retention time or returns the object unchanged if none are present.

Usage

```
applyAdjustedRtime(object)
```

Arguments

object An XCMSnExp object.

Details

Adjusted retention times are stored *in parallel* to the adjusted retention times in the XCMSnExp. The applyAdjustedRtime replaces the raw retention times (stored in the *feature data* (fData data.frame)) with the adjusted retention times.

Value

A XCMSnExp with the raw retention times being replaced with the adjusted retention time.

Note

Replacing the raw retention times with adjusted retention times disables the possibility to restore raw retention times using the dropAdjustedRtime() method. This function does **not** remove the retention time processing step with the settings of the alignment from the processHistory() of the object to ensure that the processing history is preserved.

Author(s)

Johannes Rainer

See Also

adjustRtime() for the function to perform the alignment (retention time correction).

[adjustedRtime()] for the method to extract adjusted retention times from an [XCMSnExp] object.

[dropAdjustedRtime] for the method to delete alignment results and to restore the raw retention times.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
xod <- adjustRtime(faahko_sub, param = ObiwarpParam())
hasAdjustedRtime(xod)
## Replace raw retention times with adjusted retention times.
xod <- applyAdjustedRtime(xod)
## No adjusted retention times present
hasAdjustedRtime(xod)
## Raw retention times have been replaced with adjusted retention times
plot(split(rtime(faahko_sub), fromFile(faahko_sub))[[1]] -
 split(rtime(xod), fromFile(xod))[[1]], type = "1")
## And the present history still contains the patting for the plinement
```

And the process history still contains the settings for the alignment
processHistory(xod)

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

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)</pre>
```

End(Not run)

bin, XCMSnExp-method XCMSnExp data manipulation methods inherited from MSnbase

Description

The methods listed on this page are XCMSnExp methods inherited from its parent, the OnDiskMSnExp class from the MSnbase package, that alter the raw data or are related to data subsetting. Thus calling any of these methods causes all xcms pre-processing results to be removed from the XCMSnExp object to ensure its data integrity.

bin: allows to *bin* spectra. See **bin** documentation in the MSnbase package for more details and examples.

clean: removes unused 0 intensity data points. See clean documentation in the MSnbase package for details and examples.

filterAcquisitionNum: filters the XCMSnExp object keeping only spectra with the provided acquisition numbers. See filterAcquisitionNum for details and examples.

The normalize method performs basic normalization of spectra intensities. See normalize documentation in the MSnbase package for details and examples.

The pickPeaks method performs peak picking. See pickPeaks documentation for details and examples.

The removePeaks method removes mass peaks (intensities) lower than a threshold. Note that these peaks refer to *mass* peaks, which are different to the chromatographic peaks detected and analyzed in a metabolomics experiment! See removePeaks documentation for details and examples.

The smooth method smooths spectra. See smooth documentation in MSnbase for details and examples.

Usage

```
## S4 method for signature 'XCMSnExp'
bin(x, binSize = 1L, msLevel.)
## S4 method for signature 'XCMSnExp'
clean(object, all = FALSE, verbose = FALSE, msLevel.)
## S4 method for signature 'XCMSnExp'
filterAcquisitionNum(object, n, file)
## S4 method for signature 'XCMSnExp'
normalize(object, method = c("max", "sum"), ...)
## S4 method for signature 'XCMSnExp'
pickPeaks(
   object,
   halfWindowSize = 3L,
   method = c("MAD", "SuperSmoother"),
```

```
SNR = 0L,
...
)
## S4 method for signature 'XCMSnExp'
removePeaks(object, t = "min", verbose = FALSE, msLevel.)
## S4 method for signature 'XCMSnExp'
smooth(
    x,
    method = c("SavitzkyGolay", "MovingAverage"),
    halfWindowSize = 2L,
    verbose = FALSE,
    ...
)
```

Arguments

x	XCMSnExp or OnDiskMSnExp object.
binSize	numeric(1) defining the size of a bin (in Dalton).
msLevel.	For bin, clean, filterMsLevel, removePeaks: numeric(1) defining the MS level(s) to which operations should be applied or to which the object should be subsetted.
object	XCMSnExp or OnDiskMSnExp object.
all	For clean: logical(1), if TRUE all zeros are removed.
verbose	logical(1) whether progress information should be displayed.
n	For filterAcquisitionNum: integer defining the acquisition numbers of the spectra to which the data set should be sub-setted.
file	For filterAcquisitionNum: integer defining the file index within the object to subset the object by file.
method	For normalize: character(1) specifying the normalization method. See normalize in the MSnbase package for details. For pickPeaks: character(1) defining the method. See pickPeaks for options. For smooth: character(1) defining the method. See smooth in the MSnbase package for options and details.
	Optional additional arguments.
halfWindowSize	For pickPeaks and smooth: integer(1) defining the window size for the peak picking. See pickPeaks and smooth in the MSnbase package for details and options.
SNR	For pickPeaks: numeric(1) defining the signal to noise ratio to be considered. See pickPeaks documentation for details.
t	For removePeaks: either a numeric(1) or "min" defining the threshold (method) to be used. See removePeaks for details.

Value

For all methods: a XCMSnExp object.

22

binYonX

Author(s)

Johannes Rainer

See Also

XCMSnExp-filter for methods to filter and subset XCMSnExp objects. XCMSnExp for base class documentation. OnDiskMSnExp for the documentation of the parent class.

binYonX

Aggregate values in y for bins defined on x

Description

This functions takes two same-sized numeric vectors x and y, bins/cuts x into bins (either a predefined number of equal-sized bins or bins of a pre-defined size) and aggregates values in y corresponding to x values falling within each bin. By default (i.e. method = "max") the maximal y value for the corresponding x values is identified. x is expected to be incrementally sorted and, if not, it will be internally sorted (in which case also y will be ordered according to the order of x).

Usage

```
binYonX(
  х,
  у,
  breaks,
  nBins,
  binSize,
 binFromX,
  binToX,
  fromIdx = 1L,
  toIdx = length(x),
  method = "max",
  baseValue,
  sortedX = !is.unsorted(x),
  shiftByHalfBinSize = FALSE,
  returnIndex = FALSE,
  returnX = TRUE
)
```

Arguments

х	Numeric vector to be used for binning.
У	Numeric vector (same length than x) from which the maximum values for each bin should be defined. If not provided, x will be used.
breaks	Numeric vector defining the breaks for the bins, i.e. the lower and upper values for each bin. See examples below.

nBins	integer(1) defining the number of desired bins.
binSize	numeric(1) defining the desired bin size.
binFromX	Optional numeric(1) allowing to manually specify the range of x-values to be used for binning. This will affect only the calculation of the breaks for the bins (i.e. if nBins or binSize is provided). If not provided the minimal value in the sub-set fromIdx-toIdx in input vector x will be used.
binToX	Same as binFromX, but defining the maximum x-value to be used for binning.
fromIdx	Integer vector defining the start position of one or multiple sub-sets of input vector x that should be used for binning.
toIdx	Same as toIdx, but defining the maximum index (or indices) in x to be used for binning.
method	A character string specifying the method that should be used to aggregate values in y. Allowed are "max", "min", "sum" and "mean" to identify the maximal or minimal value or to sum all values within a bin or calculate their mean value.
baseValue	The base value for empty bins (i.e. bins into which either no values in x did fall, or to which only NA values in y were assigned). By default (i.e. if not specified), NA is assigned to such bins.
sortedX	Whether x is sorted.
shiftByHalfBinS	Size
	Logical specifying whether the bins should be shifted by half the bin size to the left. Thus, the first bin will have its center at fromX and its lower and upper boundary are fromX -binSize/2 and fromX + binSize/2. This argument is ignored if breaks are provided.
returnIndex	Logical indicating whether the index of the max (if method = "max") or min (if method = "min") value within each bin in input vector x should also be reported. For methods other than "max" or "min" this argument is ignored.
returnX	<pre>logical allowing to avoid returning \$x, i.e. the mid-points of the bins. returnX = FALSE might be useful in cases where breaks are pre-defined as it consider- ably reduces the memory demand.</pre>

Details

The breaks defining the boundary of each bin can be either passed directly to the function with the argument breaks, or are calculated on the data based on arguments nBins or binSize along with fromIdx, toIdx and optionally binFromX and binToX. Arguments fromIdx and toIdx allow to specify subset(s) of the input vector x on which bins should be calculated. The default the full x vector is considered. Also, if not specified otherwise with arguments binFromX and binToX, the range of the bins within each of the sub-sets will be from x[fromIdx] to x[toIdx]. Arguments binFromX and binToX allow to overwrite this by manually defining the a range on which the breaks should be calculated. See examples below for more details.

Calculation of breaks: for nBins the breaks correspond to seq(min(x[fromIdx])),max(x[fromIdx],length.out = (nBins + 1)). For binSize the breaks correspond to seq(min(x[fromIdx]),max(x[toIdx]),by = binSize) with the exception that the last break value is forced to be equal to max(x[toIdx]). This ensures that all values from the specified range are covered by the breaks defining the bins. The last bin could however in some instances be slightly larger than binSize. See breaks_on_binSize and breaks_on_nBins for more details.

binYonX

Value

Returns a list of length 2, the first element (named "x") contains the bin mid-points, the second element (named "y") the aggregated values from input vector y within each bin. For returnIndex = TRUE the list contains an additional element "index" with the index of the max or min (depending on whether method = "max" or method = "min") value within each bin in input vector x.

Note

The function ensures that all values within the range used to define the breaks are considered in the binning (and assigned to a bin). This means that for all bins except the last one values in x have to be >= xlower and < xupper (with xlower and xupper being the lower and upper boundary, respectively). For the last bin the condition is $x \ge xlower \& x \le xupper$. Note also that if shiftByHalfBinSize is TRUE the range of values that is used for binning is expanded by binSize (i.e. the lower boundary will be fromX -binSize/2, the upper toX + binSize/2). Setting this argument to TRUE resembles the binning that is/was used in profBin function from xcms < 1.51.

NA handling: by default the function ignores NA values in y (thus inherently assumes na.rm = TRUE). No NA values are allowed in x.

Author(s)

Johannes Rainer

See Also

imputeLinInterpol

Examples

```
########
## Simple example illustrating the breaks and the binning.
##
## Define breaks for 5 bins:
brks \langle - seq(2, 12, length.out = 6)
## The first bin is then [2,4), the second [4,6) and so on.
brks
## Get the max value falling within each bin.
binYonX(x = 1:16, y = 1:16, breaks = brks)
## Thus, the largest value in x = 1:16 falling into the bin [2,4) (i.e. being
## >= 2 and < 4) is 3, the largest one falling into [4,6) is 5 and so on.
## Note however the function ensures that the minimal and maximal x-value
## (in this example 1 and 12) fall within a bin, i.e. 12 is considered for
## the last bin.
#######
## Performing the binning ons sub-set of x
##
X <- 1:16
## Bin X from element 4 to 10 into 5 bins.
X[4:10]
binYonX(X, X, nBins = 5L, fromIdx = 4, toIdx = 10)
```

```
## This defines breaks for 5 bins on the values from 4 to 10 and bins
## the values into these 5 bins. Alternatively, we could manually specify
## the range for the binning, i.e. the minimal and maximal value for the
## breaks:
binYonX(X, X, nBins = 5L, fromIdx = 4, toIdx = 10, binFromX = 1, binToX = 16)
## In this case the breaks for 5 bins were defined from a value 1 to 16 and
## the values 4 to 10 were binned based on these breaks.
#######
## Bin values within a sub-set of x, second example
##
## This example illustrates how the fromIdx and toIdx parameters can be used.
## x defines 3 times the sequence form 1 to 10, while y is the sequence from
## 1 to 30. In this very simple example x is supposed to represent M/Z values
## from 3 consecutive scans and y the intensities measured for each M/Z in
## each scan. We want to get the maximum intensities for M/Z value bins only
## for the second scan, and thus we use fromIdx = 11 and toIdx = 20. The breaks
## for the bins are defined with the nBins, binFromX and binToX.
X <- rep(1:10, 3)
Y <- 1:30
## Bin the M/Z values in the second scan into 5 bins and get the maximum
## intensity for each bin. Note that we have to specify sortedX = TRUE as
## the x and y vectors would be sorted otherwise.
binYonX(X, Y, nBins = 5L, sortedX = TRUE, fromIdx = 11, toIdx = 20)
#######
## Bin in overlapping sub-sets of X
##
## In this example we define overlapping sub-sets of X and perform the binning
## within these.
X <- 1:30
## Define the start and end indices of the sub-sets.
fIdx <- c(2, 8, 21)
tIdx <- c(10, 25, 30)
binYonX(X, nBins = 5L, fromIdx = fIdx, toIdx = tIdx)
## The same, but pre-defining also the desired range of the bins.
binYonX(X, nBins = 5L, fromIdx = fIdx, toIdx = tIdx, binFromX = 4, binToX = 28)
## The same bins are thus used for each sub-set.
```

breaks_on_binSize Generate breaks for binning using a defined bin size.

Description

Defines breaks for binSize sized bins for values ranging from fromX to toX.

Usage

breaks_on_binSize(fromX, toX, binSize)

Arguments

fromX	numeric(1) specifying the lowest value for the bins.
toX	numeric(1) specifying the largest value for the bins.
binSize	numeric(1) defining the size of a bin.

Details

This function creates breaks for bins of size binSize. The function ensures that the full data range is included in the bins, i.e. the last value (upper boundary of the last bin) is always equal toX. This however means that the size of the last bin will not always be equal to the desired bin size. See examples for more details and a comparisom to R's seq function.

Value

A numeric vector defining the lower and upper bounds of the bins.

Author(s)

Johannes Rainer

See Also

binYonX for a binning function.

Other functions to define bins: breaks_on_nBins()

Examples

```
## Define breaks with a size of 0.13 for a data range from 1 to 10:
breaks_on_binSize(1, 10, 0.13)
## The size of the last bin is however larger than 0.13:
diff(breaks_on_binSize(1, 10, 0.13))
## If we would use seq, the max value would not be included:
seq(1, 10, by = 0.13)
## In the next example we use binSize that leads to an additional last bin with
## a smaller binSize:
breaks_on_binSize(1, 10, 0.51)
## Again, the max value is included, but the size of the last bin is < 0.51.
diff(breaks_on_binSize(1, 10, 0.51))
## Using just seq would result in the following bin definition:
seq(1, 10, by = 0.51)
## Thus it defines one bin (break) less.
```

breaks_on_nBins

Description

Calculate breaks for same-sized bins for data values from fromX to toX.

Usage

```
breaks_on_nBins(fromX, toX, nBins, shiftByHalfBinSize = FALSE)
```

Arguments

fromX	numeric(1) specifying the lowest value for the bins.
toX	numeric(1) specifying the largest value for the bins.
nBins	numeric(1) defining the number of bins.
shiftByHalfBinSize	
	Logical indicating whether the bins should be shifted left by half bin size. This results centered bins, i.e. the first bin being centered at fromX and the last around toX.

Details

This generates bins such as a call to seq(fromX, toX, length.out = nBins) would. The first and second element in the result vector thus defines the lower and upper boundary for the first bin, the second and third value for the second bin and so on.

Value

A numeric vector of length nBins + 1 defining the lower and upper bounds of the bins.

Author(s)

Johannes Rainer

See Also

binYonX for a binning function.

Other functions to define bins: breaks_on_binSize()

Examples

```
## Create breaks to bin values from 3 to 20 into 20 bins
breaks_on_nBins(3, 20, nBins = 20)
## The same call but using shiftByHalfBinSize
breaks_on_nBins(3, 20, nBins = 20, shiftByHalfBinSize = TRUE)
```

c-methods

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

CalibrantMassParam-class

Calibrant mass based calibration of chromatgraphic peaks

Description

Calibrate peaks using mz values of known masses/calibrants. mz values of identified peaks are adjusted based on peaks that are close to the provided mz values. See details below for more information.

The isCalibrated function returns TRUE if chromatographic peaks of the XCMSnExp object x were calibrated and FALSE otherwise.

Usage

```
CalibrantMassParam(
  mz = list(),
  mzabs = 1e-04,
  mzppm = 5,
  neighbors = 3,
  method = "linear"
)
isCalibrated(object)
## S4 method for signature 'XCMSnExp'
```

calibrate(object, param)

Arguments

mz	a numeric or list of numeric vectors with reference mz values. If a numeric vector is provided, this is used for each sample in the XCMSnExp object. If a list is provided, it's length has to be equal to the number of samples in the experiment.
mzabs	numeric(1) the absolute error/deviation for matching peaks to calibrants (in Da).
mzppm	numeric(1) the relative error for matching peaks to calibrants in ppm (parts per million).
neighbors	integer(1) with the maximal number of peaks within the permitted distance to the calibrants that are considered. Among these the mz value of the peak with the largest intensity is used in the calibration function estimation.
method	character(1) defining the method that should be used to estimate the calibra- tion function. Can be "shift", "linear" (default) or "edgeshift".
object	An XCMSnExp object.
param	The CalibrantMassParam object with the calibration settings.

Details

The method does first identify peaks that are close to the provided mz values and, given that there difference to the calibrants is smaller than the user provided cut off (based on arguments mzabs and mzppm), their mz values are replaced with the provided mz values. The mz values of all other peaks are either globally shifted (for method = "shift" or estimated by a linear model through all calibrants. Peaks are considered close to a calibrant mz if the difference between the calibrant and its mz is <= mzabs + mz * mzppm /1e6.

Adjustment methods: adjustment function/factor is estimated using the difference between calibrant and peak mz values only for peaks that are close enough to the calibrants. The availabel methods are:

• shift: shifts the m/z of each peak by a global factor which corresponds to the average difference between peak mz and calibrant mz.

30

calibrate-methods

- linear: fits a linear model throught the differences between calibrant and peak mz values and adjusts the mz values of all peaks using this.
- edgeshift: performs same adjustment as linear for peaks that are within the mz range of the calibrants and shift outside of it.

For more information, details and examples refer to the *xcms-direct-injection* vignette.

Value

For CalibrantMassParam: a CalibrantMassParam instance. For calibrate: an XCMSnExp object with chromatographic peaks being calibrated. **Be aware** that the actual raw mz values are not (yet) calibrated, but **only** the identified chromatographic peaks.

The CalibrantMassParam function returns an instance of the CalibrantMassParam class with all settings and properties set.

The calibrate method returns an XCMSnExp object with the chromatographic peaks being calibrated. Note that **only** the detected peaks are calibrated, but not the individual mz values in each spectrum.

Note

CalibrantMassParam classes don't have exported getter or setter methods.

Author(s)

Joachim Bargsten, Johannes Rainer

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" calibrate(object, calibrants, method="linear", mzabs=0.0001, mzppm=5, neighbours=3, plo

See Also

xcmsSet-class,

chromatogram,XCMSnExp-method

Extracting chromatograms

Description

chromatogram: extract chromatographic data (such as an extracted ion chromatogram, a base peak chromatogram or total ion chromatogram) from an OnDiskMSnExp or XCMSnExp objects. See also the help page of the chromatogram function in the MSnbase package.

Usage

```
## S4 method for signature 'XCMSnExp'
chromatogram(
   object,
   rt,
   mz,
   aggregationFun = "sum",
   missing = NA_real_,
   msLevel = 1L,
   BPPARAM = bpparam(),
   adjustedRtime = hasAdjustedRtime(object),
   filled = FALSE,
   include = c("apex_within", "any", "none")
)
```

Arguments

object	Either a OnDiskMSnExp or XCMSnExp object from which the chromatograms should be extracted.
rt	numeric(2) or two-column matrix defining the lower and upper boundary for the retention time range(s). If not specified, the full retention time range of the original data will be used.
mz	numeric(2) or two-column matrix defining the lower and upper mz value for the MS data slice(s). If not specified, the chromatograms will be calculated on the full mz range.
aggregationFun	character(1) specifying the function to be used to aggregate intensity values across the mz value range for the same retention time. Allowed values are "sum" (the default), "max", "mean" and "min".
missing	numeric(1) allowing to specify the intensity value to be used if for a given re- tention time no signal was measured within the mz range of the corresponding scan. Defaults to NA_real_ (see also Details and Notes sections below). Use missing = 0 to resemble the behaviour of the getEIC from the <i>old</i> user inter- face.
msLevel	integer(1) specifying the MS level from which the chromatogram should be extracted. Defaults to msLevel = 1L.
BPPARAM	Parallelisation backend to be used, which will depend on the architecture. De-fault is BiocParallel::bparam().
adjustedRtime	For chromatogram,XCMSnExp: whether the adjusted (adjustedRtime = TRUE) or raw retention times (adjustedRtime = FALSE) should be used for filtering and returned in the resulting MChromatograms object. Adjusted retention times are used by default if available.
filled	logical(1) whether filled-in peaks should also be returned. Defaults to filled = FALSE, i.e. returns only detected chromatographic peaks in the result object.
include	character(1) defining which chromatographic peaks should be returned. Supported are include = "apex_within" (the default) which returns chromato- graphic peaks that have their apex within the mz rt range, include = "any" to return all chromatographic peaks which m/z and rt ranges overlap the mz and rt or include = "none" to not include any chromatographic peaks.

Details

Arguments rt and mz allow to specify the MS data slice (i.e. the m/z range and retention time window) from which the chromatogram should be extracted. These parameters can be either a numeric of length 2 with the lower and upper limit, or a matrix with two columns with the lower and upper limits to extract multiple EICs at once. The parameter aggregationSum allows to specify the function to be used to aggregate the intensities across the m/z range for the same retention time. Setting aggregationFun = "sum" would e.g. allow to calculate the **total ion chromatogram** (TIC), aggregationFun = "max" the **base peak chromatogram** (BPC).

If for a given retention time no intensity is measured in that spectrum a NA intensity value is returned by default. This can be changed with the parameter missing, setting missing = 0 would result in a 0 intensity being returned in these cases.

Value

chromatogram returns a XChromatograms object with the number of columns corresponding to the number of files in object and number of rows the number of specified ranges (i.e. number of rows of matrices provided with arguments mz and/or rt). All chromatographic peaks with their apex position within the m/z and retention time range are also retained as well as all feature definitions for these peaks.

Note

For XCMSnExp objects, if adjusted retention times are available, the chromatogram method will by default report and use these (for the subsetting based on the provided parameter rt). This can be changed by setting adjustedRtime = FALSE.

Author(s)

Johannes Rainer

See Also

XCMSnExp for the data object. Chromatogram for the object representing chromatographic data.

[XChromatograms] for the object allowing to arrange multiple [XChromatogram] objects.

[plot] to plot a [XChromatogram] or [MChromatograms] objects.

`as` (`as(x, "data.frame")`) in `MSnbase` for a method to extract the MS data as `data.frame`.

Examples

```
## Load a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")</pre>
```

Disable parallel processing for this example
register(SerialParam())

Extract the ion chromatogram for one chromatographic peak in the data. chrs <- chromatogram(faahko_sub, rt = c(2700, 2900), mz = 335)

chrs

```
## Identified chromatographic peaks
chromPeaks(chrs)
```

Plot the chromatogram
plot(chrs)

Extract chromatograms for multiple ranges.

34

```
mzr <- matrix(c(335, 335, 344, 344), ncol = 2, byrow = TRUE)
rtr <- matrix(c(2700, 2900, 2600, 2750), ncol = 2, byrow = TRUE)
chrs <- chromatogram(faahko_sub, mz = mzr, rt = rtr)
chromPeaks(chrs)
plot(chrs)
## Get access to all chromatograms for the second mz/rt range
chrs[1, ]
## Plot just that one
plot(chrs[1, , drop = FALSE])</pre>
```

```
chromatographic-peak-detection
```

Chromatographic peak detection methods.

Description

The findChromPeaks methods perform the chromatographic peak detection on LC/GC-MS data and are part of the modernized xcms user interface.

The implemented peak detection methods in chromatographic space are:

- **centWave** chromatographic peak detection using the *centWave* method. See centWave for more details.
- **centWave with predicted isotopes** peak detection using a two-step centWave-based approach considering also feature isotopes. See centWaveWithPredIsoROIs for more details.
- matchedFilter peak detection in chromatographic space. See matchedFilter for more details.
- **massifquant** peak detection using the Kalman filter-based method. See massifquant for more details.
- MSW single-spectrum non-chromatography MS data peak detection. See MSW for more details.

Author(s)

Johannes Rainer

See Also

findPeaks for the *old* peak detection methods.

plotChromPeaks to plot identified chromatographic peaks for one file.

highlightChromPeaks to highlight identified chromatographic peaks in an extracted ion chromatogram plot.

refineChromPeaks for methods to refine or clean identified chromatographic peaks.

manualChromPeaks to manually add/define chromatographic peaks.

Other peak detection methods: findChromPeaks-centWaveWithPredIsoROIs, findChromPeaks-centWave, findChromPeaks-massifquant, findChromPeaks-matchedFilter, findPeaks-MSW

chromPeakSpectra

Description

Extract (MS1 or MS2) spectra from an XCMSnExp object for each identified chromatographic peak. The function returns by default spectra for chromatographic peaks of **all** MS levels, but parameter peaks allows to restrict the result to selected chromatographic peaks. For msLevel = 1L (only supported for return.type = "Spectra" or return.type = "List") MS1 spectra within the retention time boundaries (in the file in which the peak was detected) are returned. For msLevel = 2L MS2 spectra are returned for a chromatographic peak if their precursor m/z is within the retention time and m/z range of the chromatographic peak. Parameter method allows to define whether all or a single spectrum should be returned:

- method = "all": (default): return all spectra for each peak.
- method = "closest_rt": return the spectrum with the retention time closest to the peak's retention time (at apex).
- method = "closest_mz": return the spectrum with the precursor m/z closest to the peaks's m/z (at apex); only supported for msLevel = 2L.
- method = "signal": return the spectrum with the sum of intensities most similar to the peak's apex signal ("maxo"); only supported for msLevel = 2L.
- method = "largest_tic": return the spectrum with the largest total signal (sum of peaks intensities).
- method = "largest_bpi": return the spectrum with the largest peak intensity (maximal peak intensity).

Parameter return.type allows to specify the *type* of the result object. Please use return.type = "Spectra" or return.type = "List", return.type = "list" or the default return.type = "MSpectra" will be deprecated (also, they do not support extracting MS1 spectra).

See also the *LC-MS/MS data analysis* vignette for more details and examples.

Usage

```
chromPeakSpectra(
    x,
    msLevel = 2L,
    expandRt = 0,
    expandMz = 0,
    ppm = 0,
    method = c("all", "closest_rt", "closest_mz", "signal", "largest_tic", "largest_bpi"),
    skipFilled = FALSE,
    return.type = c("MSpectra", "Spectra", "list", "List"),
    peaks = character()
)
```

Arguments

x	XCMSnExp object with identified chromatographic peaks.
msLevel	<pre>integer(1) defining whether MS1 or MS2 spectra should be returned. msLevel = 1 is currently only supported for return.type being "Spectra" or "List".</pre>
expandRt	numeric(1) to expand the retention time range of each peak by a constant value on each side.
expandMz	numeric(1) to expand the m/z range of each peak by a constant value on each side.
ppm	numeric(1) to expand the m/z range of each peak (on each side) by a value dependent on the peak's m/z .
method	character(1) specifying which spectra to include in the result. Defaults to method = "all". See function description for details.
skipFilled	logical(1) whether spectra for filled-in peaks should be reported or not.
return.type	<pre>character(1) defining the result type. Defaults to return.type = "MSpectra" but return.type = "Spectra" or return.type = "List" are preferred. See below for more information.</pre>
peaks	character, logical or integer allowing to specify a subset of chromato- graphic peaks in chromPeaks for which spectra should be returned (providing either their ID, a logical vector same length than nrow(chromPeaks(x)) or their index in chromPeaks(x)). This parameter overrides skipFilled and is only supported for return.type being either "Spectra" or "List".

Value

parameter return. type allow to specify the type of the returned object:

- return.type = "MSpectra": a MSpectra object with elements being Spectrum objects. The
 result objects contains all spectra for all peaks. Metadata column "peak_id" provides the ID
 of the respective peak (i.e. its rowname in chromPeaks()).
- return.type = "Spectra": a Spectra object (defined in the Spectra package). The result contains all spectra for all peaks. Metadata column "peak_id" provides the ID of the respective peak (i.e. its rowname in chromPeaks() and "peak_index" its index in the object's chromPeaks matrix.
- return.type = "list": list of lists that are either of length 0 or contain Spectrum2 object(s) within the m/z-rt range. The length of the list matches the number of peaks.
- return.type = "List": List of length equal to the number of chromatographic peaks is returned with elements being either NULL (no spectrum found) or a Spectra object.

Author(s)

Johannes Rainer

Examples

```
## Read a file with DDA LC-MS/MS data
fl <- system.file("TripleTOF-SWATH/PestMix1_DDA.mzML", package = "msdata")</pre>
dda <- readMSData(fl, mode = "onDisk")</pre>
## Subset the object to reduce runtime of the example
dda <- filterRt(dda, c(200, 400))</pre>
## Perform MS1 peak detection
dda <- findChromPeaks(dda, CentWaveParam(peakwidth = c(5, 15), prefilter = c(5, 1000)))
## Load the required Spectra package and return all MS2 spectro for each
## chromatographic peaks as a Spectra object
ms2_sps <- chromPeakSpectra(dda, return.type = "Spectra")</pre>
ms2_sps
## columns peak_id or peak_index assign spectra to the chromatographic peaks
ms2_sps$peak_id
ms2_sps$peak_index
chromPeaks(dda)
## Alternatively, return the result as a List of Spectra objects. This list
## is parallel to chromPeaks hence the mapping between chromatographic peaks
## and MS2 spectra is easier.
ms2_sps <- chromPeakSpectra(dda, return.type = "List")</pre>
ms2_sps[[1L]]
length(ms2_sps)
## In addition to MS2 spectra we could also return the MS1 spectrum for each
## chromatographic peak which is closest to the peak's apex position.
ms1_sps <- chromPeakSpectra(dda, msLevel = 1L, method = "closest_rt",</pre>
    return.type = "Spectra")
ms1_sps
## Parameter peaks would allow to extract spectra for specific peaks only
chromPeakSpectra(dda, msLevel = 1L, method = "closest_rt", peaks = c(3, 5))
```

CleanPeaksParam Remove chromatographic peaks with too large rt width

Description

Remove chromatographic peaks with a retention time range larger than the provided maximal acceptable width (maxPeakwidth).

Usage

```
CleanPeaksParam(maxPeakwidth = 10)
```

S4 method for signature 'XCMSnExp,CleanPeaksParam'
refineChromPeaks(object, param = CleanPeaksParam(), msLevel = 1L)

38

CleanPeaksParam

Arguments

maxPeakwidth	for CleanPeaksParam: numeric(1) defining the maximal allowed peak width (in retention time).
object	XCMSnExp object with identified chromatographic peaks.
param	CleanPeaksParam object defining the settings for the method.
msLevel	integer defining for which MS level(s) the chromatographic peaks should be cleaned.

Value

XCMSnExp object with chromatographic peaks exceeding the specified maximal retention time width being removed.

Note

refineChromPeaks methods will always remove feature definitions, because a call to this method can change or remove identified chromatographic peaks, which may be part of features.

Author(s)

Johannes Rainer

See Also

Other chromatographic peak refinement methods: FilterIntensityParam, MergeNeighboringPeaksParam

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")
## Disable parallel processing for this example
register(SerialParam())
## Distribution of chromatographic peak widths
quantile(chromPeaks(faahko_sub)[, "rtmax"] - chromPeaks(faahko_sub)[, "rtmin"])
## Remove all chromatographic peaks with a width larger 60 seconds
data <- refineChromPeaks(faahko_sub, param = CleanPeaksParam(60))
quantile(chromPeaks(data)[, "rtmax"] - chromPeaks(data)[, "rtmin"])
```

collect-methods

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, mzgap	, uniq 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	MSnParentPeakID	
	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	y	
	The collision energy of the fragment	

The collision energy of the fragment

Methods

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

correlate, Chromatogram, Chromatogram-method Correlate chromatograms

Description

For xcms >= 3.15.3 please use compareChromatograms() instead of correlate

Correlate intensities of two chromatograms with each other. If the two Chromatogram objects have different retention times they are first *aligned* to match data points in the first to data points in the second chromatogram. See help on alignRt in MSnbase::Chromatogram() for more details.

If correlate is called on a single MChromatograms() object a pairwise correlation of each chromatogram with each other is performed and a matrix with the correlation coefficients is returned.

Note that the correlation of two chromatograms depends also on their order, e.g. correlate(chr1, chr2) might not be identical to correlate(chr2, chr1). The lower and upper triangular part of the correlation matrix might thus be different.

Usage

```
## S4 method for signature 'Chromatogram, Chromatogram'
correlate(
 х,
 у,
 use = "pairwise.complete.obs",
 method = c("pearson", "kendall", "spearman"),
 align = c("closest", "approx"),
)
## S4 method for signature 'MChromatograms,missing'
correlate(
  х,
 y = NULL,
 use = "pairwise.complete.obs",
 method = c("pearson", "kendall", "spearman"),
 align = c("closest", "approx"),
)
## S4 method for signature 'MChromatograms, MChromatograms'
correlate(
  х,
 y = NULL,
  use = "pairwise.complete.obs",
 method = c("pearson", "kendall", "spearman"),
 align = c("closest", "approx"),
  . . .
```

)

Arguments

х	Chromatogram() or MChromatograms() object.
У	Chromatogram() or MChromatograms() object.
use	character(1) passed to the cor function. See cor() for details.
method	character(1) passed to the cor function. See cor() for details.
align	character(1) defining the alignment method to be used. See help on alignRt in MSnbase::Chromatogram() for details. The value of this parameter is passed to the method parameter of alignRt.
	optional parameters passed along to the alignRt method such as tolerance that, if set to 0 requires the retention times to be identical.

Value

numeric(1) or matrix (if called on MChromatograms objects) with the correlation coefficient. If a matrix is returned, the rows represent the chromatograms in x and the columns the chromatograms in y.

Author(s)

Michael Witting, Johannes Rainer

Examples

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.
missing	numeric(1) defining an optional value for missing values. missing = 0 would e.g. replace all NA values in the feature matrix with 0. Note that also a call to fillPeaks results in a feature matrix in which NA values are replaced by 0.
	optional arguments to be passed to mt.teststat from the multtest package.

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.

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))[2],filebase = character(),eicmax = 0,eicwidth = 200,sortpval
```

```
= TRUE,classeic = c(class1,class2),value=c("into","maxo","intb"),metlin = FALSE,h=480,w=640,mzdec=
= numeric(),...)
```

dirname

See Also

xcmsSet-class, palette

dirname

Change the file path of an OnDiskMSnExp object

Description

dirname allows to get and set the path to the directory containing the source files of the OnDiskM-SnExp (or XCMSnExp) object.

Usage

S4 method for signature 'OnDiskMSnExp'
dirname(path)

S4 replacement method for signature 'OnDiskMSnExp'
dirname(path) <- value</pre>

Arguments

path	OnDiskMSnExp.
value	character of length 1 or length equal to the number of files defining the new path to the files.

Author(s)

Johannes Rainer

do_adjustRtime_peakGroups

Align spectrum retention times across samples using peak groups found in most samples

Description

The function performs retention time correction by assessing the retention time deviation across all samples using peak groups (features) containg chromatographic peaks present in most/all samples. The retention time deviation for these features in each sample is described by fitting either a polynomial (smooth = "loess") or a linear (smooth = "linear") model to the data points. The models are subsequently used to adjust the retention time for each spectrum in each sample.

Usage

```
do_adjustRtime_peakGroups(
    peaks,
    peakIndex,
    rtime,
    minFraction = 0.9,
    extraPeaks = 1,
    smooth = c("loess", "linear"),
    span = 0.2,
    family = c("gaussian", "symmetric"),
    peakGroupsMatrix = matrix(ncol = 0, nrow = 0),
    subset = integer(),
    subsetAdjust = c("average", "previous")
)
```

Arguments

peaks	a matrix or data.frame with the identified chromatographic peaks in the samples.
peakIndex	a list of indices that provides the grouping information of the chromatographic peaks (across and within samples).
rtime	a list of numeric vectors with the retention times per file/sample.
minFraction	numeric(1) between 0 and 1 defining the minimum required fraction of samples in which peaks for the peak group were identified. Peak groups passing this criteria will aligned across samples and retention times of individual spectra will be adjusted based on this alignment. For minFraction = 1 the peak group has to contain peaks in all samples of the experiment. Note that if subset is provided, the specified fraction is relative to the defined subset of samples and not to the total number of samples within the experiment (i.e. a peak has to be present in the specified proportion of subset samples).
extraPeaks	numeric(1) defining the maximal number of additional peaks for all samples to be assigned to a peak group (i.e. feature) for retention time correction. For a data set with 6 samples, extraPeaks = 1 uses all peak groups with a total peak count <= 6 + 1. The total peak count is the total number of peaks being assigned to a peak group and considers also multiple peaks within a sample being assigned to the group.
smooth	character defining the function to be used, to interpolate corrected retention times for all peak groups. Either "loess" or "linear".
span	numeric(1) defining the degree of smoothing (if smooth = "loess"). This parameter is passed to the internal call to loess.
family	character defining the method to be used for loess smoothing. Allowed values are "gaussian" and "symmetric".See loess for more information.
peakGroupsMatrix	
	optional matrix of (raw) retention times for peak groups on which the alignment should be performed. Each column represents a sample, each row a feature/peak group. If not provided, this matrix will be determined depending on parameters

46

	minFraction and extraPeaks. If provided, minFraction and extraPeaks will be ignored.
subset	integer with the indices of samples within the experiment on which the align- ment models should be estimated. Samples not part of the subset are adjusted based on the closest subset sample. See description above for more details.
subsetAdjust	character specifying the method with which non-subset samples should be adjusted. Supported options are "previous" and "average" (default). See description above for more information.

Details

The alignment bases on the presence of compounds that can be found in all/most samples of an experiment. The retention times of individual spectra are then adjusted based on the alignment of the features corresponding to these *house keeping compounds*. The parameters minFraction and extraPeaks can be used to fine tune which features should be used for the alignment (i.e. which features most likely correspond to the above mentioned house keeping compounds).

Parameter subset allows to define a subset of samples within the experiment that should be aligned. All samples not being part of the subset will be aligned based on the adjustment of the closest sample within the subset. This allows to e.g. exclude blank samples from the alignment process with their retention times being still adjusted based on the alignment results of the *real* samples.

Value

A list with numeric vectors with the adjusted retention times grouped by sample.

Note

The method ensures that returned adjusted retention times are increasingly ordered, just as the raw retention times.

Author(s)

Colin Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" Anal. Chem. 2006, 78:779-787.

do_findChromPeaks_centWave

Core API function for centWave peak detection

Description

This function performs peak density and wavelet based chromatographic peak detection for high resolution LC/MS data in centroid mode [Tautenhahn 2008].

Usage

```
do_findChromPeaks_centWave(
 mz,
  int,
  scantime,
 valsPerSpect,
 ppm = 25,
 peakwidth = c(20, 50),
  snthresh = 10,
 prefilter = c(3, 100),
 mzCenterFun = "wMean",
  integrate = 1,
 mzdiff = -0.001,
  fitgauss = FALSE,
 noise = 0,
  verboseColumns = FALSE,
  roiList = list(),
  firstBaselineCheck = TRUE,
  roiScales = NULL,
  sleep = 0,
  extendLengthMSW = FALSE
)
```

Arguments

mz	Numeric vector with the individual m/z values from all scans/ spectra of one file/sample.
int	Numeric vector with the individual intensity values from all scans/spectra of one file/sample.
scantime	Numeric vector of length equal to the number of spectra/scans of the data repre- senting the retention time of each scan.
valsPerSpect	Numeric vector with the number of values for each spectrum.
ppm	numeric(1) defining the maximal tolerated m/z deviation in consecutive scans in parts per million (ppm) for the initial ROI definition.
peakwidth	numeric(2) with the expected approximate peak width in chromatographic space. Given as a range (min, max) in seconds.

snthresh	numeric(1) defining the signal to noise ratio cutoff.
prefilter	numeric(2): $c(k, I)$ specifying the prefilter step for the first analysis step (ROI detection). Mass traces are only retained if they contain at least k peaks with intensity >= I.
mzCenterFun	Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.
integrate	Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.
mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.
noise	numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).
verboseColumns	logical(1) whether additional peak meta data columns should be returned.
roiList	An optional list of regions-of-interest (ROI) representing detected mass traces. If ROIs are submitted the first analysis step is omitted and chromatographic peak detection is performed on the submitted ROIs. Each ROI is expected to have the following elements specified: scmin (start scan index), scmax (end scan index), mzmin (minimum m/z), mzmax (maximum m/z), length (number of scans), intensity (summed intensity). Each ROI should be represented by a list of elements or a single row data.frame.
firstBaselineC	-
	logical(1). If TRUE continuous data within regions of interest is checked to be above the first baseline.
roiScales	Optional numeric vector with length equal to roiList defining the scale for each region of interest in roiList that should be used for the centWave-wavelets.
sleep extendLengthMSN	numeric(1) defining the number of seconds to wait between iterations. Defaults to $sleep = 0$. If > 0 a plot is generated visualizing the identified chromatographic peak. Note: this argument is for backward compatibility only and will be removed in future.
	Option to force centWave to use all scales when running centWave rather than truncating with the EIC length. Uses the "open" method to extend the EIC to a integer base-2 length prior to being passed to convolve rather than the default "reflect" method. See https://github.com/sneumann/xcms/issues/445 for more information.

Details

This algorithm is most suitable for high resolution LC/{TOF,OrbiTrap,FTICR}-MS data in centroid mode. In the first phase the method identifies *regions of interest* (ROIs) representing mass traces that are characterized as regions with less than ppm m/z deviation in consecutive scans in the LC/MS map. In detail, starting with a single m/z, a ROI is extended if a m/z can be found in the next scan (spectrum) for which the difference to the mean m/z of the ROI is smaller than the user defined ppm of the m/z. The mean m/z of the ROI is then updated considering also the newly included m/z value.

These ROIs are then, after some cleanup, analyzed using continuous wavelet transform (CWT) to locate chromatographic peaks on different scales. The first analysis step is skipped, if regions of interest are passed with the roiList parameter.

Value

A matrix, each row representing an identified chromatographic peak, with columns:

mz Intensity weighted mean of m/z values of the peak across scans.

mzmin Minimum m/z of the peak.

mzmax Maximum m/z of the peak.

rt Retention time of the peak's midpoint.

rtmin Minimum retention time of the peak.

rtmax Maximum retention time of the peak.

into Integrated (original) intensity of the peak.

intb Per-peak baseline corrected integrated peak intensity.

maxo Maximum intensity of the peak.

- sn Signal to noise ratio, defined as (maxo -baseline)/sd, sd being the standard deviation of local chromatographic noise.
- egauss RMSE of Gaussian fit.

Additional columns for verboseColumns = TRUE:

mu Gaussian parameter mu.

sigma Gaussian parameter sigma.

h Gaussian parameter h.

 \mathbf{f} Region number of the m/z ROI where the peak was localized.

dppm m/z deviation of mass trace across scans in ppm.

scale Scale on which the peak was localized.

scpos Peak position found by wavelet analysis (scan number).

scmin Left peak limit found by wavelet analysis (scan number).

scmax Right peak limit found by wavelet analysis (scan numer).

50

Note

The *centWave* was designed to work on centroided mode, thus it is expected that such data is presented to the function.

This function exposes core chromatographic peak detection functionality of the *centWave* method. While this function can be called directly, users will generally call the corresponding method for the data object instead.

Author(s)

Ralf Tautenhahn, Johannes Rainer

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

centWave for the standard user interface method.

Other core peak detection functions: do_findChromPeaks_centWaveWithPredIsoROIs(), do_findChromPeaks_massifqu do_findChromPeaks_matchedFilter(), do_findPeaks_MSW()

Examples

```
## Load the test file
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")</pre>
## Subset to one file and restrict to a certain retention time range
data <- filterRt(filterFile(faahko_sub, 1), c(2500, 3000))</pre>
## Get m/z and intensity values
mzs <- mz(data)</pre>
ints <- intensity(data)</pre>
## Define the values per spectrum:
valsPerSpect <- lengths(mzs)</pre>
## Calling the function. We're using a large value for noise and prefilter
## to speed up the call in the example - in a real use case we would either
## set the value to a reasonable value or use the default value.
res <- do_findChromPeaks_centWave(mz = unlist(mzs), int = unlist(ints),</pre>
    scantime = rtime(data), valsPerSpect = valsPerSpect, noise = 10000,
    prefilter = c(3, 10000))
head(res)
```

do_findChromPeaks_centWaveWithPredIsoROIs

Core API function for two-step centWave peak detection with isotopes

Description

The do_findChromPeaks_centWaveWithPredIsoROIs performs a two-step centWave based peak detection: chromatographic peaks are identified using centWave followed by a prediction of the location of the identified peaks' isotopes in the mz-retention time space. These locations are fed as *regions of interest* (ROIs) to a subsequent centWave run. All non overlapping peaks from these two peak detection runs are reported as the final list of identified peaks.

The do_findChromPeaks_centWaveAddPredIsoROIs performs centWave based peak detection based in regions of interest (ROIs) representing predicted isotopes for the peaks submitted with argument peaks.. The function returns a matrix with the identified peaks consisting of all input peaks and peaks representing predicted isotopes of these (if found by the centWave algorithm).

Usage

```
do_findChromPeaks_centWaveWithPredIsoROIs(
  mz,
  int,
  scantime,
  valsPerSpect,
  ppm = 25,
  peakwidth = c(20, 50),
  snthresh = 10,
  prefilter = c(3, 100),
 mzCenterFun = "wMean",
  integrate = 1,
 mzdiff = -0.001,
  fitgauss = FALSE,
  noise = 0,
  verboseColumns = FALSE,
  roiList = list(),
  firstBaselineCheck = TRUE,
  roiScales = NULL,
  snthreshIsoROIs = 6.25,
 maxCharge = 3,
 maxIso = 5,
 mzIntervalExtension = TRUE,
 polarity = "unknown",
  extendLengthMSW = FALSE
)
do_findChromPeaks_addPredIsoROIs(
 mz,
```

 $do_findChromPeaks_centWaveWithPredIsoROIs$

```
int,
 scantime,
 valsPerSpect,
 ppm = 25,
 peakwidth = c(20, 50),
 snthresh = 6.25,
 prefilter = c(3, 100),
 mzCenterFun = "wMean",
 integrate = 1,
 mzdiff = -0.001,
 fitgauss = FALSE,
 noise = 0,
 verboseColumns = FALSE,
 peaks. = NULL,
 maxCharge = 3,
 maxIso = 5,
 mzIntervalExtension = TRUE,
 polarity = "unknown"
)
```

Arguments

mz	Numeric vector with the individual m/z values from all scans/ spectra of one file/sample.
int	Numeric vector with the individual intensity values from all scans/spectra of one file/sample.
scantime	Numeric vector of length equal to the number of spectra/scans of the data repre- senting the retention time of each scan.
valsPerSpect	Numeric vector with the number of values for each spectrum.
ppm	numeric(1) defining the maximal tolerated m/z deviation in consecutive scans in parts per million (ppm) for the initial ROI definition.
peakwidth	numeric(2) with the expected approximate peak width in chromatographic space. Given as a range (min, max) in seconds.
snthresh	For do_findChromPeaks_addPredIsoROIs: numeric(1) defining the signal to noise threshold for the centWave algorithm. For do_findChromPeaks_centWaveWithPredIsoROIs: numeric(1) defining the signal to noise threshold for the initial (first) centWave run.
prefilter	numeric(2): $c(k, I)$ specifying the prefilter step for the first analysis step (ROI detection). Mass traces are only retained if they contain at least k peaks with intensity >= I.
mzCenterFun	Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.

	integrate	Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.
	mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
	fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.
	noise	numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).
	verboseColumns	logical(1) whether additional peak meta data columns should be returned.
	roiList	An optional list of regions-of-interest (ROI) representing detected mass traces. If ROIs are submitted the first analysis step is omitted and chromatographic peak detection is performed on the submitted ROIs. Each ROI is expected to have the following elements specified: scmin (start scan index), scmax (end scan index), mzmin (minimum m/z), mzmax (maximum m/z), length (number of scans), intensity (summed intensity). Each ROI should be represented by a list of elements or a single row data.frame.
	firstBaselineCh	neck
		logical(1). If TRUE continuous data within regions of interest is checked to be above the first baseline.
	roiScales	Optional numeric vector with length equal to roiList defining the scale for each region of interest in roiList that should be used for the centWave-wavelets.
	snthreshIsoROIs	
		numeric(1) defining the signal to noise ratio cutoff to be used in the second centWave run to identify peaks for predicted isotope ROIs.
	maxCharge	<pre>integer(1) defining the maximal isotope charge. Isotopes will be defined for charges 1:maxCharge.</pre>
	maxIso	integer(1) defining the number of isotope peaks that should be predicted for each peak identified in the first centWave run.
mzIntervalExtension		
		logical(1) whether the mz range for the predicted isotope ROIs should be extended to increase detection of low intensity peaks.
	polarity	character(1) specifying the polarity of the data. Currently not used, but has to be "positive", "negative" or "unknown" if provided.
	extendLengthMSW	V
		Option to force centWave to use all scales when running centWave rather than truncating with the EIC length. Uses the "open" method to extend the EIC to a integer base-2 length prior to being passed to convolve rather than the default "reflect" method. See https://github.com/sneumann/xcms/issues/445 for more information.

do_findChromPeaks_centWaveWithPredIsoROIs

peaks. A matrix or xcmsPeaks object such as one returned by a call to link{do_findChromPeaks_centWave} or link{findPeaks.centWave} (both with verboseColumns = TRUE) with the peaks for which isotopes should be predicted and used for an additional peak detectoin using the centWave method. Required columns are: "mz", "mzmin", "mzmax", "scmin", "scmax", "scale" and "into".

Details

For more details on the centWave algorithm see centWave.

Value

A matrix, each row representing an identified chromatographic peak. All non-overlapping peaks identified in both centWave runs are reported. The matrix columns are:

mz Intensity weighted mean of m/z values of the peaks across scans.

mzmin Minimum m/z of the peaks.

mzmax Maximum m/z of the peaks.

rt Retention time of the peak's midpoint.

rtmin Minimum retention time of the peak.

rtmax Maximum retention time of the peak.

into Integrated (original) intensity of the peak.

intb Per-peak baseline corrected integrated peak intensity.

maxo Maximum intensity of the peak.

sn Signal to noise ratio, defined as (maxo -baseline)/sd, sd being the standard deviation of local chromatographic noise.

egauss RMSE of Gaussian fit.

Additional columns for verboseColumns = TRUE:

mu Gaussian parameter mu.

sigma Gaussian parameter sigma.

h Gaussian parameter h.

f Region number of the m/z ROI where the peak was localized.

dppm m/z deviation of mass trace across scans in ppm.

scale Scale on which the peak was localized.

scpos Peak position found by wavelet analysis (scan number).

scmin Left peak limit found by wavelet analysis (scan number).

scmax Right peak limit found by wavelet analysis (scan numer).

Author(s)

Hendrik Treutler, Johannes Rainer

See Also

Other core peak detection functions: do_findChromPeaks_centWave(), do_findChromPeaks_massifquant(), do_findChromPeaks_matchedFilter(), do_findPeaks_MSW()

do_findChromPeaks_massifquant

Core API function for massifquant peak detection

Description

Massifquant is a Kalman filter (KF)-based chromatographic peak detection for XC-MS data in centroid mode. The identified peaks can be further refined with the *centWave* method (see do_findChromPeaks_centWave for details on centWave) by specifying withWave = TRUE.

Usage

```
do_findChromPeaks_massifquant(
 mz,
  int,
  scantime,
  valsPerSpect,
  ppm = 10,
 peakwidth = c(20, 50),
  snthresh = 10,
 prefilter = c(3, 100),
 mzCenterFun = "wMean",
  integrate = 1,
 mzdiff = -0.001,
  fitgauss = FALSE,
 noise = 0,
  verboseColumns = FALSE,
  criticalValue = 1.125,
  consecMissedLimit = 2,
  unions = 1,
  checkBack = 0,
 withWave = FALSE
```

)

Arguments

Numeric vector with the individual m/z values from all scans/ spectra of one file/sample.
Numeric vector with the individual intensity values from all scans/spectra of one file/sample.
Numeric vector of length equal to the number of spectra/scans of the data repre- senting the retention time of each scan.

valsPerSpect	Numeric vector with the number of values for each spectrum.	
ppm	numeric(1) defining the maximal tolerated m/z deviation in consecutive scans in parts per million (ppm) for the initial ROI definition.	
peakwidth	numeric(2) with the expected approximate peak width in chromatographic space. Given as a range (min, max) in seconds.	
snthresh	numeric(1) defining the signal to noise ratio cutoff.	
prefilter	numeric(2): c(k, I) specifying the prefilter step for the first analysis step (ROI detection). Mass traces are only retained if they contain at least k peaks with intensity >= I.	
mzCenterFun	Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.	
integrate	Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.	
mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.	
fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.	
noise	numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).	
verboseColumns	logical(1) whether additional peak meta data columns should be returned.	
criticalValue	numeric(1). 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 peak 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 peak in the XC-MS sample have a small mass deviance in ppm error, a smaller critical value might be better and vice versa.	
consecMissedLimit		
	integer(1) Suggested values: (1,2,3). While a peak 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 peak.	
unions	integer (1) set to 1 if apply t-test union on segmentation; set to \emptyset if no t-test to be applied on chromatographically continous peaks sharing same m/z range. Explanation: With very few data points, sometimes a Kalman Filter stops tracking a peak prematurely. Another Kalman Filter is instantiated and begins following	

	the rest of the signal. Because tracking is done backwards to forwards, this algo- rithmic defect leaves a real peak divided into two segments or more. With this option turned on, the program identifies segmented peaks and combines them (merges them) into one with a two sample t-test. The potential danger of this option is that some truly distinct peaks may be merged.
checkBack	integer(1) set to 1 if turned on; set to 0 if turned off. The convergence of a Kalman Filter to a peak's precise m/z mapping is very fast, but sometimes it incorporates erroneous centroids as part of a peak (especially early on). The scanBack 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 peak 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.
withWave	logical(1) if TRUE, the peaks identified first with Massifquant are subsequently filtered with the second step of the centWave algorithm, which includes wavelet estimation.

Details

This algorithm's performance has been tested rigorously on high resolution LC/OrbiTrap, TOF-MS data in centroid mode. Simultaneous kalman filters identify peaks 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 peak 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

A matrix, each row representing an identified chromatographic peak, with columns:

mz Intensity weighted mean of m/z values of the peaks across scans.

mzmin Minumum m/z of the peak.

mzmax Maximum m/z of the peak.

rtmin Minimum retention time of the peak.

rtmax Maximum retention time of the peak.

rt Retention time of the peak's midpoint.

into Integrated (original) intensity of the peak.

maxo Maximum intensity of the peak.

If withWave is set to TRUE, the result is the same as returned by the do_findChromPeaks_centWave method.

Author(s)

Christopher Conley

References

Conley CJ, Smith R, Torgrip RJ, Taylor RM, Tautenhahn R and Prince JT "Massifquant: opensource Kalman filter-based XC-MS isotope trace feature detection" *Bioinformatics* 2014, 30(18):2636-43.

See Also

massifquant for the standard user interface method.

Other core peak detection functions: do_findChromPeaks_centWaveWithPredIsoROIs(), do_findChromPeaks_centWave do_findChromPeaks_matchedFilter(), do_findPeaks_MSW()

Examples

```
## Load the test file
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")</pre>
## Subset to one file and restrict to a certain retention time range
data <- filterRt(filterFile(faahko_sub, 1), c(2500, 3000))</pre>
## Get m/z and intensity values
mzs <- mz(data)</pre>
ints <- intensity(data)</pre>
## Define the values per spectrum:
valsPerSpect <- lengths(mzs)</pre>
## Perform the peak detection using massifquant - setting prefilter to
## a high value to speed up the call for the example
res <- do_findChromPeaks_massifquant(mz = unlist(mzs), int = unlist(ints),</pre>
    scantime = rtime(data), valsPerSpect = valsPerSpect,
    prefilter = c(3, 10000)
head(res)
```

Description

This function identifies peaks in the chromatographic time domain as described in [Smith 2006]. The intensity values are binned by cutting The LC/MS data into slices (bins) of a mass unit (binSize

m/z) wide. Within each bin the maximal intensity is selected. The peak detection is then performed in each bin by extending it based on the steps parameter to generate slices comprising bins current_bin -steps +1 to current_bin + steps -1. Each of these slices is then filtered with matched filtration using a second-derative Gaussian as the model peak shape. After filtration peaks are detected using a signal-to-ration cut-off. For more details and illustrations see [Smith 2006].

Usage

```
do_findChromPeaks_matchedFilter(
 mz,
  int,
  scantime,
  valsPerSpect,
 binSize = 0.1,
  impute = "none",
  baseValue,
  distance,
  fwhm = 30,
  sigma = fwhm/2.3548,
 max = 5,
  snthresh = 10,
  steps = 2,
 mzdiff = 0.8 - binSize * steps,
  index = FALSE,
  sleep = 0
)
```

Arguments

mz	Numeric vector with the individual m/z values from all scans/ spectra of one file/sample.
int	Numeric vector with the individual intensity values from all scans/spectra of one file/sample.
scantime	Numeric vector of length equal to the number of spectra/scans of the data repre- senting the retention time of each scan.
valsPerSpect	Numeric vector with the number of values for each spectrum.
binSize	numeric(1) specifying the width of the bins/slices in m/z dimension.
impute	Character string specifying the method to be used for missing value imputation. Allowed values are "none" (no linear interpolation), "lin" (linear interpola- tion), "linbase" (linear interpolation within a certain bin-neighborhood) and "intlin". See imputeLinInterpol for more details.
baseValue	The base value to which empty elements should be set. This is only considered for method = "linbase" and corresponds to the profBinLinBase's baselevel argument.
distance	For method = "linbase": number of non-empty neighboring element of an empty element that should be considered for linear interpolation. See details section for more information.

fwhm	numeric(1) specifying the full width at half maximum of matched filtration gaussian model peak. Only used to calculate the actual sigma, see below.
sigma	numeric(1) specifying the standard deviation (width) of the matched filtration model peak.
max	numeric(1) representing the maximum number of peaks that are expected/will be identified per slice.
snthresh	numeric(1) defining the signal to noise ratio cutoff.
steps	numeric(1) defining the number of bins to be merged before filtration (i.e. the number of neighboring bins that will be joined to the slice in which filtration and peak detection will be performed).
mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
index	logical(1) specifying whether indicies should be returned instead of values for m/z and retention times.
sleep	numeric(1) defining the number of seconds to wait between iterations. Defaults to $sleep = 0$. If > 0 a plot is generated visualizing the identified chromatographic peak. Note: this argument is for backward compatibility only and will be removed in future.

Details

The intensities are binned by the provided m/z values within each spectrum (scan). Binning is performed such that the bins are centered around the m/z values (i.e. the first bin includes all m/z values between min(mz) -bin_size/2 and min(mz) + bin_size/2).

For more details on binning and missing value imputation see binYonX and imputeLinInterpol methods.

Value

A matrix, each row representing an identified chromatographic peak, with columns:

mz Intensity weighted mean of m/z values of the peak across scans.

mzmin Minimum m/z of the peak.

mzmax Maximum m/z of the peak.

rt Retention time of the peak's midpoint.

rtmin Minimum retention time of the peak.

rtmax Maximum retention time of the peak.

into Integrated (original) intensity of the peak.

intf Integrated intensity of the filtered peak.

maxo Maximum intensity of the peak.

maxf Maximum intensity of the filtered peak.

i Rank of peak in merged EIC (<= max).

sn Signal to noise ratio of the peak

This function exposes core peak detection functionality of the *matchedFilter* method. While this function can be called directly, users will generally call the corresponding method for the data object instead (e.g. the link{findPeaks.matchedFilter} method).

Author(s)

Colin A Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" *Anal. Chem.* 2006, 78:779-787.

See Also

binYonX for a binning function, imputeLinInterpol for the interpolation of missing values. matchedFilter for the standard user interface method.

Other core peak detection functions: do_findChromPeaks_centWaveWithPredIsoROIs(), do_findChromPeaks_centWave do_findChromPeaks_massifquant(), do_findPeaks_MSW()

Examples

62

do_findPeaks_MSW

Description

This function performs peak detection in mass spectrometry direct injection spectrum using a wavelet based algorithm.

Usage

```
do_findPeaks_MSW(mz, int, snthresh = 3, verboseColumns = FALSE, ...)
```

Arguments

mz	Numeric vector with the individual m/z values from all scans/ spectra of one file/sample.
int	Numeric vector with the individual intensity values from all scans/spectra of one file/sample.
snthresh	numeric(1) defining the signal to noise ratio cutoff.
verboseColumns	logical(1) whether additional peak meta data columns should be returned.
	Additional parameters to be passed to the peakDetectionCWT function.

Details

This is a wrapper around the peak picker in Bioconductor's MassSpecWavelet package calling peakDetectionCWT and tuneInPeakInfo functions. See the *xcmsDirect* vignette for more information.

Value

A matrix, each row representing an identified peak, with columns:

mz m/z value of the peak at the centroid position.

mzmin Minimum m/z of the peak.

mzmax Maximum m/z of the peak.

rt Always -1.

rtmin Always -1.

rtmax Always -1.

into Integrated (original) intensity of the peak.

maxo Maximum intensity of the peak.

intf Always NA.

maxf Maximum MSW-filter response of the peak.

sn Signal to noise ratio.

Author(s)

Joachim Kutzera, Steffen Neumann, Johannes Rainer

See Also

MSW for the standard user interface method. peakDetectionCWT from the MassSpecWavelet package.

Other core peak detection functions: do_findChromPeaks_centWaveWithPredIsoROIs(), do_findChromPeaks_centWave do_findChromPeaks_massifquant(), do_findChromPeaks_matchedFilter()

```
do_groupChromPeaks_density
```

Core API function for peak density based chromatographic peak grouping

Description

The do_groupChromPeaks_density function performs chromatographic peak grouping based on the density (distribution) of peaks, found in different samples, along the retention time axis in slices of overlapping mz ranges.

Usage

```
do_groupChromPeaks_density(
   peaks,
   sampleGroups,
   bw = 30,
   minFraction = 0.5,
   minSamples = 1,
   binSize = 0.25,
   maxFeatures = 50,
   sleep = 0
)
```

Arguments

peaks	A matrix or data.frame with the mz values and retention times of the identi- fied chromatographic peaks in all samples of an experiment. Required columns are "mz", "rt" and "sample". The latter should contain numeric values repre- senting the index of the sample in which the peak was found.
sampleGroups	A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the same group).

bw	numeric(1) defining the bandwidth (standard deviation ot the smoothing ker- nel) to be used. This argument is passed to the [density() method.
minFraction	numeric(1) defining the minimum fraction of samples in at least one sample group in which the peaks have to be present to be considered as a peak group (feature).
minSamples	numeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).
binSize	numeric(1) defining the size of the overlapping slices in mz dimension.
maxFeatures	numeric(1) with the maximum number of peak groups to be identified in a single mz slice.
sleep	numeric(1) defining the time to <i>sleep</i> between iterations and plot the result from the current iteration.

Details

For overlapping slices along the mz dimension, the function calculates the density distribution of identified peaks along the retention time axis and groups peaks from the same or different samples that are close to each other. See (Smith 2006) for more details.

Value

A data.frame, each row representing a (mz-rt) feature (i.e. a peak group) with columns:

- "mzmed": median of the peaks' apex mz values.
- "mzmin": smallest mz value of all peaks' apex within the feature.
- "mzmax":largest mz value of all peaks' apex within the feature.
- "rtmed": the median of the peaks' retention times.
- "rtmin": the smallest retention time of the peaks in the group.
- "rtmax": the largest retention time of the peaks in the group.
- "npeaks": the total number of peaks assigned to the feature.
- "peakidx": a list with the indices of all peaks in a feature in the peaks input matrix.

Note that this number can be larger than the total number of samples, since multiple peaks from the same sample could be assigned to a feature.

Note

The default settings might not be appropriate for all LC/GC-MS setups, especially the bw and binSize parameter should be adjusted accordingly.

Author(s)

Colin Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" Anal. Chem. 2006, 78:779-787.

See Also

Other core peak grouping algorithms: do_groupChromPeaks_nearest(), do_groupPeaks_mzClust()

Examples

```
## Load the test file
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Extract the matrix with the identified peaks from the xcmsSet:
pks <- chromPeaks(faahko_sub)
## Perform the peak grouping with default settings:
res <- do_groupChromPeaks_density(pks, sampleGroups = rep(1, 3))
## The feature definitions:
head(res)
```

 $do_groupChromPeaks_nearest$

Core API function for chromatic peak grouping using a nearest neighbor approach

Description

The do_groupChromPeaks_nearest function groups peaks across samples by creating a master peak list and assigning corresponding peaks from all samples to each peak group (i.e. feature). The method is inspired by the correspondence algorithm of mzMine (Katajamaa 2006).

Usage

```
do_groupChromPeaks_nearest(
    peaks,
    sampleGroups,
    mzVsRtBalance = 10,
    absMz = 0.2,
    absRt = 15,
    kNN = 10
)
```

Arguments

peaks	A matrix or data.frame with the mz values and retention times of the identi- fied chromatographic peaks in all samples of an experiment. Required columns are "mz", "rt" and "sample". The latter should contain numeric values repre- senting the index of the sample in which the peak was found.
sampleGroups	A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the same group).
mzVsRtBalance	numeric(1) representing the factor by which mz values are multiplied before calculating the (euclician) distance between two peaks.
absMz	numeric(1) maximum tolerated distance for mz values.
absRt	numeric(1) maximum tolerated distance for rt values.
kNN	numeric(1) representing the number of nearest neighbors to check.

Value

A list with elements "featureDefinitions" and "peakIndex". "featureDefinitions" is a matrix, each row representing an (mz-rt) feature (i.e. peak group) with columns:

- "mzmed": median of the peaks' apex mz values.
- "mzmin": smallest mz value of all peaks' apex within the feature.
- "mzmax":largest mz value of all peaks' apex within the feature.
- "rtmed": the median of the peaks' retention times.
- "rtmin": the smallest retention time of the peaks in the feature.
- "rtmax": the largest retention time of the peaks in the feature.
- "npeaks": the total number of peaks assigned to the feature.

"peakIndex" is a list with the indices of all peaks in a feature in the peaks input matrix.

References

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

See Also

Other core peak grouping algorithms: do_groupChromPeaks_density(), do_groupPeaks_mzClust()

do_groupPeaks_mzClust Core API function for peak grouping using mzClust

Description

The do_groupPeaks_mzClust function performs high resolution correspondence on single spectra samples.

Usage

```
do_groupPeaks_mzClust(
    peaks,
    sampleGroups,
    ppm = 20,
    absMz = 0,
    minFraction = 0.5,
    minSamples = 1
)
```

Arguments

peaks	A matrix or data.frame with the mz values and retention times of the identi- fied chromatographic peaks in all samples of an experiment. Required columns are "mz", "rt" and "sample". The latter should contain numeric values repre- senting the index of the sample in which the peak was found.
sampleGroups	A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the same group).
ppm	numeric(1) representing the relative mz error for the clustering/grouping (in parts per million).
absMz	numeric(1) representing the absolute mz error for the clustering.
minFraction	numeric(1) defining the minimum fraction of samples in at least one sample group in which the peaks have to be present to be considered as a peak group (feature).
minSamples	numeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).

Value

A list with elements "featureDefinitions" and "peakIndex". "featureDefinitions" is a matrix, each row representing an (mz-rt) feature (i.e. peak group) with columns:

- "mzmed": median of the peaks' apex mz values.
- "mzmin": smallest mz value of all peaks' apex within the feature.

- "mzmax": largest mz value of all peaks' apex within the feature.
- "rtmed": always -1.
- "rtmin": always -1.
- "rtmax": always -1.
- "npeaks": the total number of peaks assigned to the feature. Note that this number can be larger than the total number of samples, since multiple peaks from the same sample could be assigned to a group.

"peakIndex" is a list with the indices of all peaks in a peak group in the peaks input matrix.

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

Other core peak grouping algorithms: do_groupChromPeaks_density(), do_groupChromPeaks_nearest()

estimatePrecursorIntensity

Estimate precursor intensity for MS level 2 spectra

Description

estimatePrecursorIntensity determines the precursor intensity for a MS 2 spectrum based on the intensity of the respective signal from the neighboring MS 1 spectra (i.e. based on the peak with the m/z matching the precursor m/z of the MS 2 spectrum). Based on parameter method either the intensity of the peak from the previous MS 1 scan is used (method = "previous") or an interpolation between the intensity from the previous and subsequent MS1 scan is used (method = "interpolation", which considers also the retention times of the two MS1 scans and the retention time of the MS2 spectrum).

Usage

```
estimatePrecursorIntensity(
    x,
    ppm = 10,
    method = c("previous", "interpolation"),
    BPPARAM = bpparam()
)
```

Arguments

х	OnDiskMSnExp or XCMSnExp object.
ppm	numeric(1) defining the maximal acceptable difference (in ppm) of the precursor m/z and the m/z of the corresponding peak in the MS 1 scan.
method	character(1) defining the method how the precursor intensity should be deter- mined (see description above for details). Defaults to method = "previous".
BPPARAM	parallel processing setup. See <pre>bpparam()</pre> for details.

Value

numeric with length equal to the number of spectra in x. NA is returned for MS 1 spectra or if no matching peak in a MS 1 scan can be found for an MS 2 spectrum

Author(s)

Johannes Rainer

etg

Empirically Transformed Gaussian function

Description

A general function for asymmetric chromatographic peaks.

Usage

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

Arguments

times to evaluate function at
peak height
time of leading edge inflection point
time of trailing edge inflection point
leading edge parameter
trailing edge parameter
leading edge parameter
trailing edge parameter
leading edge parameter
trailing edge parameter

Value

The function evaluated at times x.

exportMetaboAnalyst

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

exportMetaboAnalyst Export data for use in MetaboAnalyst

Description

Export the feature table for further analysis in the MetaboAnalyst software (or the MetaboAnalystR R package).

Usage

```
exportMetaboAnalyst(
    x,
    file = NULL,
    label,
    value = "into",
    digits = NULL,
    groupnames = FALSE,
    ...
)
```

Arguments

Х	XCMSnExp object with identified chromatographic peaks grouped across samples.
file	character(1) defining the file name. If not specified, the matrix with the content is returned.
label	either character(1) specifying the phenodata column in x defining the sample grouping or a vector with the same length than samples in x defining the group assignment of the samples.
value	character(1) specifying the value to be returned for each feature. See featureValues() for more details.
digits	integer(1) defining the number of significant digits to be used for numeric. The default NULL uses getOption("digits"). See format() for more infor- mation.

groupnames	logical(1) whether row names of the resulting matrix should be the feature
	IDs (groupnames = FALSE; default) or IDs that are composed of the m/z and re-
	tention time of the features (in the format M <m z="">T<rt> (groupnames = TRUE).</rt></m>
	See help of the groupnames function for details.
	additional parameters to be passed to the featureValues() function.

Value

If file is not specified, the function returns the matrix in the format supported by MetaboAnalyst.

Author(s)

Johannes Rainer

extractMsData,OnDiskMSnExp-method DEPRECATED: Extract a data.frame containing MS data

Description

UPDATE: the extractMsData and plotMsData functions are deprecated and as(x, "data.frame") and plot(x, type = "XIC") (x being an OnDiskMSnExp or XCMSnExp object) should be used instead. See examples below. Be aware that filtering the raw object might however drop the adjusted retention times. In such cases it is advisable to use the applyAdjustedRtime() function prior to filtering.

Extract a data.frame of retention time, mz and intensity values from each file/sample in the provided rt-mz range (or for the full data range if rt and mz are not defined).

Usage

```
## S4 method for signature 'OnDiskMSnExp'
extractMsData(object, rt, mz, msLevel = 1L)
## S4 method for signature 'XCMSnExp'
extractMsData(
   object,
   rt,
   mz,
   msLevel = 1L,
   adjustedRtime = hasAdjustedRtime(object)
)
```

Arguments

object	A XCMSnExp or OnDiskMSnExp object.
rt	numeric(2) with the retention time range from which the data should be extracted.
mz	numeric(2) with the mz range.
msLevel	integer defining the MS level(s) to which the data should be sub-setted prior to extraction; defaults to $msLevel = 1L$.
adjustedRtime	(for extractMsData,XCMSnExp): logical(1) specifying if adjusted or raw re- tention times should be reported. Defaults to adjusted retention times, if these are present in object.

Value

A list of length equal to the number of samples/files in object. Each element being a data.frame with columns "rt", "mz" and "i" with the retention time, mz and intensity tuples of a file. If no data is available for the mz-rt range in a file a data.frame with 0 rows is returned for that file.

Author(s)

Johannes Rainer

See Also

XCMSnExp for the data object.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")
## Disable parallel processing for this example
register(SerialParam())
## Extract the full MS data for a certain retention time range
## as a data.frame
tmp <- filterRt(faahko_sub, rt = c(2800, 2900))
ms_all <- as(tmp, "data.frame")
head(ms_all)
nrow(ms_all)
```

feature-grouping

Description

Feature *compounding* aims at identifying and grouping LC-MS features representing different ions or adducts (including isotopes) of the same originating compound. The MsFeatures package provides a general framework and functionality to group features based on different properties. The groupFeatures methods for XCMSnExp objects implemented in xcms extend these to enable the *compounding* of LC-MS data. Note that these functions simply define feature groups but don't actually *aggregate* or combine the features.

See MsFeatures::groupFeatures() for an overview on the general feature grouping concept as well as details on the individual settings and parameters.

The available options for groupFeatures on xcms preprocessing results (i.e. on XCMSnExp objects after correspondence analysis with groupChromPeaks()) are:

- Grouping by similar retention times: groupFeatures-similar-rtime().
- Grouping by similar feature values across samples: AbundanceSimilarityParam().
- Grouping by similar peak shape of extracted ion chromatograms: EicSimilarityParam().

An ideal workflow grouping features should sequentially perform the above methods (in the listed order).

Compounded feature groups can be accessed with the featureGroups function.

Usage

```
## S4 method for signature 'XCMSnExp'
featureGroups(object)
```

```
## S4 replacement method for signature 'XCMSnExp'
featureGroups(object) <- value</pre>
```

Arguments

object	an XCMSnExp() object.
value	for featureGroups<-: replacement for the feature groups in object. Has to be of length 1 or length equal to the number of features in object.

Author(s)

Johannes Rainer, Mar Garcia-Aloy, Vinicius Veri Hernandes

See Also

plotFeatureGroups() for visualization of grouped features.

featureChromatograms Extract ion chromatograms for each feature

Description

Extract ion chromatograms for features in an XCMSnExp object. The function returns for each feature its extracted ion chromatogram and all associated peaks with it. The chromatogram is extracted from the m/z - rt region including all chromatographic peaks of that features (i.e. based on the ranges of "mzmin", "mzmax", "rtmin", "rtmax" of all chromatographic peaks of the feature).

By default only chromatographic peaks associated with a feature are included for an extracted ion chromatogram (parameter include = "feature_only"). By setting include = "apex_within" all chromatographic peaks (and eventually the feature which they are part of - if feature definitions are present) that have their apex position within the m/z - rt range from which the chromatographic peaks (and eventually e "any" or include = "all" all chromatographic peaks (and eventually the feature between the e and the chromatographic peaks are returned too. With include = "any" or include = "all" all chromatographic peaks (and eventually the feature in which they are present) overlapping the m/z and rt range will be returned.

Usage

```
featureChromatograms(
    x,
    expandRt = 0,
    aggregationFun = "max",
    features,
    include = c("feature_only", "apex_within", "any", "all"),
    filled = FALSE,
    n = length(fileNames(x)),
    value = c("maxo", "into"),
    ...
)
```

Arguments

x	XCMSnExp object with grouped chromatographic peaks.
expandRt	numeric(1) to expand the retention time range for each chromatographic peak by a constant value on each side.
aggregationFun	character(1) specifying the name that should be used to aggregate intensity values across the m/z value range for the same retention time. The default "sum" returns a base peak chromatogram.
features	integer, character or logical defining a subset of features for which chro- matograms should be returned. Can be the index of the features in featureDefinitions, feature IDs (row names of featureDefinitions) or a logical vector.
include	character(1) defining which chromatographic peaks (and related feature def- initions) should be included in the returned XChromatograms(). Defaults to "feature_only"; See description above for options and details.

filled	logical(1) whether filled-in peaks should be included in the result object. The default is filled = FALSE, i.e. only detected peaks are reported.
n	integer(1) to optionally specify the number of $top n$ samples from which the EIC should be extracted.
value	character(1) specifying the column to be used to sort the samples. Can be either "maxo" (the default) or "into" to use the maximal peak intensity or the integrated peak area, respectively.
	optional arguments to be passed along to the chromatogram() function.

Value

XChromatograms() object.

Note

When extracting EICs from only the top n samples it can happen that one or more of the features specified with features are dropped because they have no detected peak in the *top n* samples. The chance for this to happen is smaller if x contains also filled-in peaks (with fillChromPeaks).

Author(s)

Johannes Rainer

See Also

filterColumnsKeepTop() to filter the extracted EICs keeping only the *top n* columns (samples) with the highest intensity.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Subset the object to a smaller retention time range
xdata <- filterRt(faahko_sub, c(2500, 3500))
xdata <- groupChromPeaks(xdata,
    param = PeakDensityParam(minFraction = 0.8, sampleGroups = rep(1, 3)))
## Get the feature definitions
featureDefinitions(xdata)
## Extract ion chromatograms for the first 3 features. Parameter
## `features` can be either the feature IDs or feature indices.
chrs <- featureChromatograms(xdata, features = 1:3)</pre>
```

```
## Plot the XIC for the first feature using different colors for each file
plot(chrs[1, ], col = c("red", "green", "blue"))
```

featureSpectra

Extract spectra associated with features

Description

This function returns spectra associated with the identified features in the input object. By default, spectra are returned for all features (from all MS levels), but parameter features allows to specify selected features for which the result should be returned. Parameter msLevel allows to define whether MS level 1 or 2 spectra should be returned. For msLevel = 1L all MS1 spectra within the retention time range of each chromatographic peak (in that respective data file) associated with a feature are returned. Note that for samples in which no peak was identified (or even filled-in) no spectra are returned. For msLevel = 2L all MS2 spectra with a retention time within the retention time range and their precursor m/z within the m/z range of any chromatographic peak of a feature are returned. See also chromPeakSpectra() (used internally to extract spectra for each chromatographic peak of a feature) for additional information.

In contrast to the chromPeakSpectra() function, selecting a method different than "all" will not return a single spectrum per feature, but one spectrum per chromatographic peak assigned to the feature.

Note also that msLevel = 1L is only supported for return.type = "List" or return.type = "Spectra".

Usage

```
featureSpectra(
    x,
    msLevel = 2L,
    expandRt = 0,
    expandMz = 0,
    ppm = 0,
    skipFilled = FALSE,
    return.type = c("MSpectra", "Spectra", "list", "List"),
    features = character(),
    ...
)
```

Arguments

х	XCMSnExp object with feature defitions available.
msLevel	<pre>integer(1) defining whether MS1 or MS2 spectra should be returned. msLevel = 1 is currently only supported for return.type being "Spectra" or "List".</pre>
expandRt	numeric(1) to expand the retention time range of each peak by a constant value on each side.

expandMz	numeric(1) to expand the m/z range of each peak by a constant value on each side.
ppm	numeric(1) to expand the m/z range of each peak (on each side) by a value dependent on the peak's m/z.
skipFilled	logical(1) whether spectra for filled-in peaks should be reported or not.
return.type	<pre>character(1) defining the result type. Defaults to return.type = "MSpectra" but return.type = "Spectra" or return.type = "List" are preferred. See below for more information.</pre>
features	character, logical or integer allowing to specify a subset of features in featureDefinitions for which spectra should be returned (providing either their ID, a logical vector same length than nrow(featureDefinitions(x)) or their index in featureDefinitions(x)). This parameter overrides skipFilled and is only supported for return.type being either "Spectra" or "List".
• • •	additional arguments to be passed along to chromPeakSpectra(), such as method.

Value

parameter return. type allow to specify the type of the returned object:

- return.type = "MSpectra": a MSpectra object with elements being Spectrum objects. The result objects contains all spectra for all features. Metadata column "feature_id" provides the ID of the respective feature (i.e. its rowname in featureDefinitions()).
- return.type = "Spectra": a Spectra object (defined in the Spectra package). The result contains all spectra for all features. Metadata column "feature_id" provides the ID of the respective feature (i.e. its rowname in featureDefinitions().
- return.type = "list": list of lists that are either of length 0 or contain Spectrum2 object(s) within the m/z-rt range. The length of the list matches the number of features.
- return.type = "List": List of length equal to the number of features with MS level msLevel is returned with elements being either NULL (no spectrum found) or a Spectra object.

Author(s)

Johannes Rainer

featureSummary Simple feature summaries

Description

Simple function to calculate feature summaries. These include counts and percentages of samples in which a chromatographic peak is present for each feature and counts and percentages of samples in which more than one chromatographic peak was annotated to the feature. Also relative standard deviations (RSD) are calculated for the integrated peak areas per feature across samples. For 'perSampleCounts = TRUE' also the individual chromatographic peak counts per sample are returned.

featureSummary

Usage

```
featureSummary(
    x,
    group,
    perSampleCounts = FALSE,
    method = "maxint",
    skipFilled = TRUE
)
```

Arguments

x	'XCMSnExp' object with correspondence results.
group	'numeric', 'logical', 'character' or 'factor' with the same length than 'x' has samples to aggregate counts by the groups defined in 'group'.
perSampleCount	S
	'logical(1)' whether feature wise individual peak counts per sample should be returned too.
method	'character' passed to the [featureValues()] function. See respective help page for more information.
skipFilled	'logical(1)' whether filled-in peaks should be excluded (default) or included in the summary calculation.

Value

'matrix' with one row per feature and columns:

- '"count"': the total number of samples in which a peak was found. - '"perc"': the percentage of samples in which a peak was found. - '"multi_count"': the total number of samples in which more than one peak was assigned to the feature. - '"multi_perc"': the percentage of those samples in which a peak was found, that have also multiple peaks annotated to the feature. Example: for a feature, at least one peak was detected in 50 samples. In 5 of them 2 peaks were assigned to the feature. '"multi_perc"' is in this case 10 - '"rsd"': relative standard deviation (coefficient of variation) of the integrated peak area of the feature's peaks. - The same 4 columns are repeated for each unique element (level) in 'group' if 'group' was provided.

If 'perSampleCounts = TRUE' also one column for each sample is returned with the peak counts per sample.

Author(s)

Johannes Rainer

FillChromPeaksParam-class

Integrate areas of missing peaks

Description

expandMz,expandMz<-: getter and setter for the expandMz slot of the object.

expandRt,expandRt<-: getter and setter for the expandRt slot of the object.

ppm,ppm<-: getter and setter for the ppm slot of the object.

Integrate signal in the mz-rt area of a feature (chromatographic peak group) for samples in which no chromatographic peak for this feature was identified and add it to the chromPeaks() matrix. Such *filled-in* peaks are indicated with a TRUE in column "is_filled" in the result object's chromPeakData() data frame.

Two different gap-filling approaches are implemented:

- param = FillChromPeaksParam(): the default of the original xcms code. Signal is integrated from the m/z and retention time range as defined in the featureDefinitions() data frame, i.e. from the "rtmin", "rtmax", "mzmin" and "mzmax". See details below for more information and settings for this method.
- param = ChromPeakAreaParam(): the area from which the signal for a feature is integrated is defined based on the feature's chromatographic peak areas. The m/z range is by default defined as the the lower quartile of chromatographic peaks' "mzmin" value to the upper quartile of the chromatographic peaks' "mzmax" values. The retention time range for the area is defined analogously. Alternatively, by setting mzmin = median, mzmax = median, rtmin = median and rtmax = median in ChromPeakAreaParam, the median "mzmin", "mzmax", "rtmin" and "rtmax" values from all detected chromatographic peaks of a feature would be used instead. In contrast to the FillChromPeaksParam approach this method uses the actual identified chromatographic peaks of a feature to define the area from which the signal should be integrated.

Usage

```
FillChromPeaksParam(
    expandMz = 0,
    expandRt = 0,
    ppm = 0,
    fixedMz = 0,
    fixedRt = 0
)
fixedRt(object)
fixedMz(object)
ChromPeakAreaParam(
```

```
mzmin = function(z) quantile(z, probs = 0.25),
```

```
mzmax = function(z) quantile(z, probs = 0.75),
 rtmin = function(z) quantile(z, probs = 0.25),
  rtmax = function(z) quantile(z, probs = 0.75)
)
## S4 method for signature 'FillChromPeaksParam'
expandMz(object)
## S4 replacement method for signature 'FillChromPeaksParam'
expandMz(object) <- value</pre>
## S4 method for signature 'FillChromPeaksParam'
expandRt(object)
## S4 replacement method for signature 'FillChromPeaksParam'
expandRt(object) <- value</pre>
## S4 method for signature 'FillChromPeaksParam'
ppm(object)
## S4 replacement method for signature 'FillChromPeaksParam'
ppm(object) <- value</pre>
## S4 method for signature 'XCMSnExp,FillChromPeaksParam'
fillChromPeaks(object, param, msLevel = 1L, BPPARAM = bpparam())
## S4 method for signature 'XCMSnExp,ChromPeakAreaParam'
fillChromPeaks(object, param, msLevel = 1L, BPPARAM = bpparam())
```

S4 method for signature 'XCMSnExp,missing'
fillChromPeaks(object, param, BPPARAM = bpparam(), msLevel = 1L)

Arguments

expandMz	for FillChromPeaksParam: numeric(1) defining the value by which the mz width of peaks should be expanded. Each peak is expanded in mz direction by expandMz * their original m/z width. A value of 0 means no expansion, a value of 1 grows each peak by 1 * the m/z width of the peak resulting in peaks with twice their original size in m/z direction (expansion by half m/z width to both sides).
expandRt	for FillChromPeaksParam: numeric(1), same as expandMz but for the reten- tion time width.
ppm	for FillChromPeaksParam: numeric(1) optionally specifying a <i>ppm</i> by which the m/z width of the peak region should be expanded. For peaks with an m/z width smaller than mean(c(mzmin,mzmax)) * ppm / 1e6, the mzmin will be re- placed by mean(c(mzmin,mzmax)) - (mean(c(mzmin,mzmax)) * ppm / 2 / 1e6) mzmax by mean(c(mzmin,mzmax)) + (mean(c(mzmin,mzmax)) * ppm / 2 / 1e6) This is applied before eventually expanding the m/z width using the expandMz

	parameter.
fixedMz	for FillChromPeaksParam: numeric(1) defining a constant factor by which the m/z width of each feature is to be expanded. The m/z width is expanded on both sides by fixedMz (i.e. fixedMz is subtracted from the lower m/z and added to the upper m/z). This expansion is applied <i>after</i> expandMz and ppm.
fixedRt	for FillChromPeaksParam: numeric(1) defining a constant factor by which the retention time width of each factor is to be expanded. The rt width is expanded on both sides by fixedRt (i.e. fixedRt is subtracted from the lower rt and added to the upper rt). This expansion is applied <i>after</i> expandRt.
object	XCMSnExp object with identified and grouped chromatographic peaks.
mzmin	function to be applied to values in the "mzmin" column of all chromatographic peaks of a feature to define the lower m/z value of the area from which signal for the feature should be integrated. Defaults to mzmin = function(z) quantile(z,probs = 0.25) hence using the 25% quantile of all values.
mzmax	function to be applied to values in the "mzmax" column of all chromatographic peaks of a feature to define the upper m/z value of the area from which signal for the feature should be integrated. Defaults to mzmax = function(z) quantile(z,probs = 0.75) hence using the 75% quantile of all values.
rtmin	function to be applied to values in the "rtmin" column of all chromatographic peaks of a feature to define the lower rt value of the area from which signal for the feature should be integrated. Defaults to rtmin = function(z) quantile(z,probs = 0.25) hence using the 25% quantile of all values.
rtmax	function to be applied to values in the "rtmax" column of all chromatographic peaks of a feature to define the upper rt value of the area from which signal for the feature should be integrated. Defaults to rtmax = function(z) quantile(z,probs = 0.75) hence using the 75% quantile of all values.
value	The value for the slot.
param	FillChromPeaksParam or ChromPeakAreaParam object defining which approach should be used (see details section).
msLevel	integer(1) defining the MS level on which peak filling should be performed (defaults to msLevel = 1L). Only peak filling on one MS level at a time is supported, to fill in peaks for MS level 1 and 2 run first using msLevel = 1 and then (on the returned result object) again with msLevel = 2.
BPPARAM	Parallel processing settings.

Details

After correspondence (i.e. grouping of chromatographic peaks across samples) there will always be features (peak groups) that do not include peaks from every sample. The fillChromPeaks method defines intensity values for such features in the missing samples by integrating the signal in the mz-rt region of the feature. Two different approaches to define this region are available: with ChromPeakAreaParam the region is defined based on the detected **chromatographic peaks** of a feature, while with FillChromPeaksParam the region is defined based on the m/z and retention times of the **feature** (which represent the m/z and retention times of the apex position of the associated chromatographic peaks). For the latter approach various parameters are available to

FillChromPeaksParam-class

increase the area from which signal is to be integrated, either by a constant value (fixedMz and fixedRt) or by a feature-relative amount (expandMz and expandRt).

Adjusted retention times will be used if available.

Based on the peak finding algorithm that was used to identify the (chromatographic) peaks, different internal functions are used to guarantee that the integrated peak signal matches as much as possible the peak signal integration used during the peak detection. For peaks identified with the matchedFilter() method, signal integration is performed on the *profile matrix* generated with the same settings used also during peak finding (using the same bin size for example). For direct injection data and peaks identified with the MSW algorithm signal is integrated only along the mz dimension. For all other methods the complete (raw) signal within the area is used.

Value

The FillChromPeaksParam function returns a FillChromPeaksParam object.

A XCMSnExp object with previously missing chromatographic peaks for features filled into its chromPeaks() matrix.

Slots

expandMz, expandRt, ppm, fixedMz, fixedRt See corresponding parameter above.

rtmin, rtmax, mzmin, mzmax See corresponding parameter above.

Note

The reported "mzmin", "mzmax", "rtmin" and "rtmax" for the filled peaks represents the actual MS area from which the signal was integrated. Note that no peak is filled in if no signal was present in a file/sample in the respective mz-rt area. These samples will still show a NA in the matrix returned by the featureValues() method.

Author(s)

Johannes Rainer

See Also

groupChromPeaks() for methods to perform the correspondence.

featureArea for the function to define the m/z-retention time region for each feature.

Examples

```
## Load a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
res <- faahko_sub
## Disable parallel processing for this example
register(SerialParam())
```

```
## Perform the correspondence. We assign all samples to the same group.
res <- groupChromPeaks(res,</pre>
    param = PeakDensityParam(sampleGroups = rep(1, length(fileNames(res)))))
## For how many features do we lack an integrated peak signal?
sum(is.na(featureValues(res)))
## Filling missing peak data using the peak area from identified
## chromatographic peaks.
res <- fillChromPeaks(res, param = ChromPeakAreaParam())</pre>
## How many missing values do we have after peak filling?
sum(is.na(featureValues(res)))
## Get the peaks that have been filled in:
fp <- chromPeaks(res)[chromPeakData(res)$is_filled, ]</pre>
head(fp)
## Get the process history step along with the parameters used to perform
## The peak filling:
ph <- processHistory(res, type = "Missing peak filling")[[1]]</pre>
ph
## The parameter class:
ph@param
## It is also possible to remove filled-in peaks:
res <- dropFilledChromPeaks(res)</pre>
sum(is.na(featureValues(res)))
```

fillPeaks-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
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

fillPeaks.chrom-methods

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	(DEPRECATED): number of slaves/cores to be used for parallel peak filling. 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. Users are advised to use the BPPARAM parameter instead.
expand.mz	Expansion factor for the m/z range used for integration.
expand.rt	Expansion factor for the rentention time range used for integration.
BPPARAM	allows to define a specific parallel processing setup for the current task (see bpparam from the BiocParallel package help more information). The default uses the globally defined parallel setup.

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

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

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)

Note

In contrast to the fillPeaks.chrom method the maximum intensity reported in column "maxo" is not the maximum intensity measured in the expected peak area (defined by columns "mzmin" and "mzmax"), but the largest intensity of mz value(s) closest to the "mzmed" of the feature.

See Also

xcmsSet-class, getPeaks fillPeaks

filterColumnsIntensityAbove,MChromatograms-method Filtering sets of chromatographic data

Description

These functions allow to filter (subset) MChromatograms() or XChromatograms() objects, i.e. sets of chromatographic data, without changing the data (intensity and retention times) within the individual chromatograms (Chromatogram() objects).

- filterColumnsIntensityAbove: subsets a MChromatograms objects keeping only columns (samples) for which value is larger than the provided threshold in which rows (i.e. if which = "any" a column is kept if **any** of the chromatograms in that column have a value larger than threshold or with which = "all" **all** chromatograms in that column fulfill this criteria). Parameter value allows to define on which value the comparison should be performed, with value = "bpi" the maximum intensity of each chromatogram is compared to threshold, with value = "tic" the total sum of intensities of each chromatogram is compared to threshold. For XChromatogramsobject, value = "maxo" and value = "into" are supported which compares the largest intensity of all identified chromatographic peaks in the chromatogram with-threshold', or the integrated peak area, respectively.
- filterColumnsKeepTop: subsets a MChromatograms object keeping the top n columns sorted by the value specified with sortBy. In detail, for each column the value defined by sortBy is extracted from each chromatogram and aggregated using the aggregationFun. Thus, by default, for each chromatogram the maximum intensity is determined (sortBy = "bpi") and these values are summed up for chromatograms in the same column (aggregationFun = sum). The columns are then sorted by these values and the top n columns are retained in the returned MChromatograms. Similar to the filterColumnsIntensityAbove function, this function allows to use for XChromatograms objects to sort the columns by column sortBy = "maxo" or sortBy = "into" of the chromPeaks matrix.

Usage

```
## S4 method for signature 'MChromatograms'
filterColumnsIntensityAbove(
   object,
   threshold = 0,
   value = c("bpi", "tic"),
```

```
which = c("any", "all")
)
## S4 method for signature 'MChromatograms'
filterColumnsKeepTop(
 object,
 n = 1L,
 sortBy = c("bpi", "tic"),
  aggregationFun = sum
)
## S4 method for signature 'XChromatograms'
filterColumnsIntensityAbove(
 object,
  threshold = 0,
 value = c("bpi", "tic", "maxo", "into"),
which = c("any", "all")
)
## S4 method for signature 'XChromatograms'
filterColumnsKeepTop(
 object,
 n = 1L,
 sortBy = c("bpi", "tic", "maxo", "into"),
 aggregationFun = sum
)
```

Arguments

object	<pre>MChromatograms() or XChromatograms() object.</pre>
threshold	for filterColumnsIntensityAbove: numeric(1) with the threshold value to compare against.
value	<pre>character(1) defining which value should be used in the comparison or sort- ing. Can be value = "bpi" (default) to use the maximum intensity per chro- matogram or value = "tic" to use the sum of intensities per chromatogram. For XChromatograms() objects also value = "maxo" and value = "into" is supported to use the maximum intensity or the integrated area of identified chro- matographic peaks in each chromatogram.</pre>
which	for filterColumnsIntensityAbove: character(1) defining whether any (which = "any", default) or all (which = "all") chromatograms in a column have to fulfill the criteria for the column to be kept.
n	for filterColumnsKeepTop: integer(1) specifying the number of columns that should be returned. n will be rounded to the closest (larger) integer value.
sortBy	for filterColumnsKeepTop: the value by which columns should be ordered to determine the top n columns. Can be either sortBy = "bpi" (the default), in which case the maximum intensity of each column's chromatograms is used, or sortBy = "tic" to use the total intensity sum of all chromatograms. For

XChromatograms() objects also value = "maxo" and value = "into" is supported to use the maximum intensity or the integrated area of identified chromatographic peaks in each chromatogram.

aggregationFun for filterColumnsKeepTop: function to be used to aggregate (combine) the values from all chromatograms in each column. Defaults to aggregationFun = sum in which case the sum of the values is used to rank the columns. Alternatively the mean, median or similar function can be used.

Value

a filtered MChromatograms (or XChromatograms) object with the same number of rows (EICs) but eventually a lower number of columns (samples).

Author(s)

Johannes Rainer

Examples

```
chr1 <- Chromatogram(rtime = 1:10 + rnorm(n = 10, sd = 0.3),
    intensity = c(5, 29, 50, NA, 100, 12, 3, 4, 1, 3))
chr2 <- Chromatogram(rtime = 1:10 + rnorm(n = 10, sd = 0.3),
    intensity = c(80, 50, 20, 10, 9, 4, 3, 4, 1, 3))
chr3 <- Chromatogram(rtime = 3:9 + rnorm(7, sd = 0.3),
    intensity = c(53, 80, 130, 15, 5, 3, 2))
chrs <- MChromatograms(list(chr1, chr2, chr1, chr3, chr2, chr3),</pre>
    ncol = 3, byrow = FALSE)
chrs
#### filterColumnsIntensityAbove
##
## Keep all columns with for which the maximum intensity of any of its
## chromatograms is larger 90
filterColumnsIntensityAbove(chrs, threshold = 90)
## Require that ALL chromatograms in a column have a value larger 90
filterColumnsIntensityAbove(chrs, threshold = 90, which = "all")
## If none of the columns fulfills the criteria no columns are returned
filterColumnsIntensityAbove(chrs, threshold = 900)
## Filtering XChromatograms allow in addition to filter on the columns
## "maxo" or "into" of the identified chromatographic peaks within each
## chromatogram.
#### filterColumnsKeepTop
##
## Keep the 2 columns with the highest sum of maximal intensities in their
## chromatograms
filterColumnsKeepTop(chrs, n = 1)
```

```
## Keep the 50 percent of columns with the highest total sum of signal. Note
## that n will be rounded to the next larger integer value
filterColumnsKeepTop(chrs, n = 0.5 * ncol(chrs), sortBy = "tic")
```

filterFeatureDefinitions

XCMSnExp filtering and subsetting

Description

The methods listed on this page allow to filter and subset XCMSnExp objects. Most of them are inherited from the OnDiskMSnExp object defined in the MSnbase package and have been adapted for XCMSnExp to enable correct subsetting of preprocessing results.

- [: subset a XCMSnExp object by spectra. Be aware that this removes **all** preprocessing results, except adjusted retention times if keepAdjustedRtime = TRUE is passed to the method.
- [[: extracts a single Spectrum object (defined in MSnbase). The reported retention time is the adjusted retention time if alignment has been performed.
- filterChromPeaks: subset the chromPeaks matrix in object. Parameter method allows to specify how the chromatographic peaks should be filtered. Currently, only method = "keep" is supported which allows to specify chromatographic peaks to keep with parameter keep (i.e. provide a logical, integer or character defining which chromatographic peaks to keep). Feature definitions (if present) are updated correspondingly.
- filterFeatureDefinitions: allows to subset the feature definitions of an XCMSnExp object. Parameter features allow to define which features to keep. It can be a logical, integer (index of features to keep) or character (feature IDs) vector.
- filterFile: allows to reduce the XCMSnExp to data from only selected files. Identified chromatographic peaks for these files are retained while correspondence results (feature definitions) are removed by default. To force keeping feature definitions use keepFeatures = TRUE. Adjusted retention times (if present) are retained by default if present. Use keepAdjustedRtime = FALSE to drop them.
- filterMsLevel: reduces the XCMSnExp object to spectra of the specified MS level(s). Chromatographic peaks and identified features are also subsetted to the respective MS level. See also the filterMsLevel documentation in MSnbase for details and examples.
- filterMz: filters the data set based on the provided m/z value range. All chromatographic peaks and features (grouped peaks) falling **completely** within the provided mz value range are retained (i.e. if their minimal m/z value is >= mz[1] and the maximal m/z value <= mz[2]. Adjusted retention times, if present, are kept.
- filterRt: filters the data set based on the provided retention time range. All chromatographic peaks and features (grouped peaks) **completely** within the specified retention time window are retained (i.e. if the retention time corresponding to the peak's apex is within the specified rt range). If retention time correction has been performed, the method will by default filter the object by adjusted retention times. The argument adjusted allows to specify manually whether filtering should be performed on raw or adjusted retention times. Filtering by retention time does not drop any preprocessing results nor does it remove or change alignment results (i.e. adjusted retention times). The method returns an empty object if no spectrum or feature is within the specified retention time range.

• split: splits an XCMSnExp object into a list of XCMSnExp objects based on the provided parameter f. Note that by default all pre-processing results are removed by the splitting, except adjusted retention times, if the optional argument keepAdjustedRtime = TRUE is provided.

Usage

```
filterFeatureDefinitions(x, features)
## S4 method for signature 'XCMSnExp,ANY,ANY,ANY'
x[i, j, ..., drop = TRUE]
## S4 method for signature 'XCMSnExp,ANY,ANY'
x[[i, j, drop = FALSE]]
## S4 method for signature 'XCMSnExp'
filterMsLevel(object, msLevel., keepAdjustedRtime = hasAdjustedRtime(object))
## S4 method for signature 'XCMSnExp'
filterFile(
  object,
  file,
  keepAdjustedRtime = hasAdjustedRtime(object),
  keepFeatures = FALSE
)
## S4 method for signature 'XCMSnExp'
filterMz(object, mz, msLevel., ...)
## S4 method for signature 'XCMSnExp'
filterRt(object, rt, msLevel., adjusted = hasAdjustedRtime(object))
## S4 method for signature 'XCMSnExp,ANY'
split(x, f, drop = FALSE, ...)
## S4 method for signature 'XCMSnExp'
filterChromPeaks(
  object,
  keep = rep(TRUE, nrow(chromPeaks(object))),
  method = "keep",
  . . .
)
```

Arguments

x For [and [[: an XCMSnExp object.
 features For filterFeatureDefinitions: either a integer specifying the indices of the features (rows) to keep, a logical with a length matching the number of rows of featureDefinitions or a character with the feature (row) names.

i	For [: numeric or logical vector specifying to which spectra the data set should be reduced. For [[: a single integer or character.
j	For [and [[: not supported.
	Optional additional arguments.
drop	For [and [[: not supported.
object	A XCMSnExp object.
msLevel.	For filterMz, filterRt: numeric defining the MS level(s) to which operations should be applied or to which the object should be subsetted.
keepAdjustedRt	
	For filterFile, filterMsLevel, [, split: logical(1) defining whether the adjusted retention times should be kept, even if e.g. features are being removed (and the retention time correction was performed on these features).
file	For filterFile: integer defining the file index within the object to subset the object by file or character specifying the file names to sub set. The indices are expected to be increasingly ordered, if not they are ordered internally.
keepFeatures	For filterFile: logical(1) whether correspondence results (feature defini- tions) should be kept or dropped. Defaults to keepFeatures = FALSE hence feature definitions are removed from the returned object by default.
mz	For filterMz: numeric(2) defining the lower and upper mz value for the filtering.
rt	For filterRt: numeric(2) defining the retention time window (lower and upper bound) for the filtering.
adjusted	For filterRt: logical indicating whether the object should be filtered by original (adjusted = FALSE) or adjusted retention times (adjusted = TRUE). For spectra: whether the retention times in the individual Spectrum objects should be the adjusted or raw retention times.
f	For split a vector of length equal to the length of x defining how x should be splitted. It is converted internally to a factor.
keep	For filterChromPeaks: logical, integer or character defining which chro- matographic peaks should be retained.
method	For filterChromPeaks: character(1) allowing to specify the method by which chromatographic peaks should be filtered. Currently only method = "keep" is supported (i.e. specify with parameter keep which chromatographic peaks should be retained).

Details

All subsetting methods try to ensure that the returned data is consistent. Correspondence results for example are removed by default if the data set is sub-setted by file, since the correspondence results are dependent on the files on which correspondence was performed. This can be changed by setting keepFeatures = TRUE. For adjusted retention times, most subsetting methods support the argument keepAdjustedRtime (even the [method) that forces the adjusted retention times to be retained even if the default would be to drop them.

Value

All methods return an XCMSnExp object.

Note

The filterFile method removes also process history steps not related to the files to which the object should be sub-setted and updates the fileIndex attribute accordingly. Also, the method does not allow arbitrary ordering of the files or re-ordering of the files within the object.

Note also that most of the filtering methods, and also the subsetting operations [drop all or selected preprocessing results. To consolidate the alignment results, i.e. ensure that adjusted retention times are always preserved, use the applyAdjustedRtime() function on the object that contains the alignment results. This replaces the raw retention times with the adjusted ones.

Author(s)

Johannes Rainer

See Also

XCMSnExp for base class documentation.

XChromatograms() for similar filter functions on XChromatograms objects.

Examples

```
## Loading a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")</pre>
## Disable parallel processing for this example
register(SerialParam())
## Subset the dataset to the first and third file.
xod_sub <- filterFile(faahko_sub, file = c(1, 3))</pre>
## The number of chromatographic peaks per file for the full object
table(chromPeaks(faahko_sub)[, "sample"])
## The number of chromatographic peaks per file for the subset
table(chromPeaks(xod_sub)[, "sample"])
basename(fileNames(faahko_sub))
basename(fileNames(xod_sub))
## Filter on mz values; chromatographic peaks and features within the
## mz range are retained (as well as adjusted retention times).
xod_sub <- filterMz(faahko_sub, mz = c(300, 400))</pre>
head(chromPeaks(xod_sub))
nrow(chromPeaks(xod_sub))
nrow(chromPeaks(faahko_sub))
```

```
## Filter on rt values. All chromatographic peaks and features within the
## retention time range are retained. Filtering is performed by default on
## adjusted retention times, if present.
xod_sub <- filterRt(faahko_sub, rt = c(2700, 2900))</pre>
range(rtime(xod_sub))
head(chromPeaks(xod_sub))
range(chromPeaks(xod_sub)[, "rt"])
nrow(chromPeaks(faahko_sub))
nrow(chromPeaks(xod_sub))
## Extract a single Spectrum
faahko_sub[[4]]
## Subsetting using [ removes all preprocessing results - using
## keepAdjustedRtime = TRUE would keep adjusted retention times, if present.
xod_sub <- faahko_sub[fromFile(faahko_sub) == 1]</pre>
xod_sub
## Using split does also remove preprocessing results, but it supports the
## optional parameter keepAdjustedRtime.
## Split the object into a list of XCMSnExp objects, one per file
xod_list <- split(faahko_sub, f = fromFile(faahko_sub))</pre>
xod_list
```

FilterIntensityParam Remove chromatographic peaks based on intensity

Description

Remove chromatographic peaks with intensities below the specified threshold. By default, with nValues = 1, all peaks with an intensity >= threshold are retained. Parameter value allows to specify the column of the chromPeaks() matrix that should be used for the filtering (defaults to value = "maxo" and thus evaluating the maximal intensity for each peak). With nValues > 1 it is possible to keep only peaks that have nValues intensities >= threshold. Note that this requires data import from the original MS files and run time of the call can thus be significantly larger. Also, for nValues > 1 parameter value is ignored.

Usage

```
FilterIntensityParam(threshold = 0, nValues = 1L, value = "maxo")
## S4 method for signature 'XCMSnExp,FilterIntensityParam'
refineChromPeaks(
   object,
   param = FilterIntensityParam(),
   msLevel = 1L,
```

BPPARAM = bpparam()

Arguments

)

threshold	numeric(1) defining the minimal required intensity for a peak to be retained. Defaults to threshold = 0.
nValues	integer(1) defining the number of data points (per chromatographic peak) that have to be \geq threshold. Defaults to nValues = 1.
value	character(1) specifying the column in chromPeaks() that should be used for the comparison. This is ignored for nValues > 1.
object	XCMSnExp object with identified chromatographic peaks.
param	FilterIntensityParam object defining the settings for the method.
msLevel	integer(1) defining the MS level in which peaks should be filtered.
BPPARAM	parameter object to set up parallel processing. Uses the default parallel process- ing setup returned by bpparam(). See bpparam() for details and examples.

Value

XCMSnExp object with filtererd chromatographic peaks.

Author(s)

Johannes Rainer, Mar Garcia-Aloy

See Also

Other chromatographic peak refinement methods: CleanPeaksParam, MergeNeighboringPeaksParam

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Remove all peaks with a maximal intensity below 50000
res <- refineChromPeaks(faahko_sub, param = FilterIntensityParam(threshold = 50000))
nrow(chromPeaks(faahko_sub))
nrow(chromPeaks(res))
all(chromPeaks(res)[, "maxo"] > 50000)
## Keep only chromatographic peaks that have 3 signals above 20000; we
## perform this on the data of a single file.
```

```
xdata <- filterFile(faahko_sub)</pre>
```

```
res <- refineChromPeaks(xdata, FilterIntensityParam(threshold = 20000, nValues = 3))
nrow(chromPeaks(xdata))
nrow(chromPeaks(res))</pre>
```

Description

findChromPeaks on a Chromatogram or MChromatograms object with a CentWaveParam parameter object performs centWave-based peak detection on purely chromatographic data. See centWave for details on the method and CentWaveParam for details on the parameter class. Note that not all settings from the CentWaveParam will be used. See peaksWithCentWave() for the arguments used for peak detection on purely chromatographic data.

After chromatographic peak detection, identified peaks can also be *refined* with the refineChromPeaks() method, which can help to reduce peak detection artifacts.

Usage

S4 method for signature 'Chromatogram,CentWaveParam'
findChromPeaks(object, param, ...)

S4 method for signature 'MChromatograms,CentWaveParam'
findChromPeaks(object, param, BPPARAM = bpparam(), ...)

S4 method for signature 'MChromatograms,MatchedFilterParam'
findChromPeaks(object, param, BPPARAM = BPPARAM, ...)

Arguments

object	a Chromatogram or MChromatograms object.
param	a CentWaveParam object specifying the settings for the peak detection. See peaksWithCentWave() for the description of arguments used for peak detection.
	currently ignored.
BPPARAM	a parameter class specifying if and how parallel processing should be performed (only for XChromatograms objects). It defaults to bpparam(). See bpparam() for more information.

Value

If called on a Chromatogram object, the method returns an XChromatogram object with the identified peaks. See peaksWithCentWave() for details on the peak matrix content.

Author(s)

Johannes Rainer

See Also

peaksWithCentWave() for the downstream function and centWave for details on the method.

Examples

```
## Loading a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")</pre>
faahko_sub <- filterRt(faahko_sub, c(2500, 3700))</pre>
##
od <- as(filterFile(faahko_sub, 1L), "OnDiskMSnExp")</pre>
## Extract chromatographic data for a small m/z range
chr <- chromatogram(od, mz = c(272.1, 272.3))[1, 1]
## Identify peaks with default settings
xchr <- findChromPeaks(chr, CentWaveParam())</pre>
xchr
## Plot data and identified peaks.
plot(xchr)
## Perform peak detection on an MChromatograms object
od3 <- readMSData(c(system.file("cdf/KO/ko15.CDF", package = "faahKO"),</pre>
    system.file("cdf/KO/ko16.CDF", package = "faahKO"),
    system.file("cdf/KO/ko18.CDF", package = "faahKO")),
    mode = "onDisk")
## Disable parallel processing for this example
register(SerialParam())
## Extract chromatograms for a m/z - retention time slice
chrs <- chromatogram(od3, mz = 344, rt = c(2500, 3500))
## Perform peak detection using CentWave
xchrs <- findChromPeaks(chrs, param = CentWaveParam())</pre>
xchrs
## Extract the identified chromatographic peaks
chromPeaks(xchrs)
## plot the result
plot(xchrs)
```

Description

findChromPeaks on a Chromatogram or MChromatograms object with a MatchedFilterParam parameter object performs matchedFilter-based peak detection on purely chromatographic data. See matchedFilter for details on the method and MatchedFilterParam for details on the parameter class. Note that not all settings from the MatchedFilterParam will be used. See peaksWithMatchedFilter() for the arguments used for peak detection on purely chromatographic data.

Usage

S4 method for signature 'Chromatogram,MatchedFilterParam'
findChromPeaks(object, param, ...)

Arguments

object	a Chromatogram or MChromatograms object.
param	a MatchedFilterParam object specifying the settings for the peak detection. See peaksWithMatchedFilter() for the description of arguments used for peak detection.
	currently ignored.

Value

If called on a Chromatogram object, the method returns a matrix with the identified peaks. See peaksWithMatchedFilter() for details on the matrix content.

Author(s)

Johannes Rainer

See Also

peaksWithMatchedFilter() for the downstream function and matchedFilter for details on the method.

Examples

```
## Loading a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")
faahko_sub <- filterRt(faahko_sub, c(2500, 3700))</pre>
```

##

findChromPeaks-centWave

```
od <- as(filterFile(faahko_sub, 1L), "OnDiskMSnExp")
## Extract chromatographic data for a small m/z range
chr <- chromatogram(od, mz = c(272.1, 272.3))[1, 1]
## Identify peaks with default settings
xchr <- findChromPeaks(chr, MatchedFilterParam())
## Plot the identified peaks
plot(xchr)</pre>
```

findChromPeaks-centWave

Chromatographic peak detection using the centWave method

Description

The centWave algorithm perform peak density and wavelet based chromatographic peak detection for high resolution LC/MS data in centroid mode [Tautenhahn 2008].

The CentWaveParam class allows to specify all settings for a chromatographic peak detection using the centWave method. Instances should be created with the CentWaveParam constructor.

The detectChromPeaks, OnDiskMSnExp, CentWaveParam method performs chromatographic peak detection using the *centWave* algorithm on all samples from an OnDiskMSnExp object. OnDiskMSnExp objects encapsule all experiment specific data and load the spectra data (mz and intensity values) on the fly from the original files applying also all eventual data manipulations.

ppm,ppm<-: getter and setter for the ppm slot of the object.

peakwidth,peakwidth<-: getter and setter for the peakwidth slot of the object.

snthresh, snthresh<-: getter and setter for the snthresh slot of the object.

prefilter, prefilter <-: getter and setter for the prefilter slot of the object.

mzCenterFun,mzCenterFun<-: getter and setter for the mzCenterFun slot of the object.

integrate, integrate <-: getter and setter for the integrate slot of the object.

mzdiff,mzdiff<-: getter and setter for the mzdiff slot of the object.

fitgauss, fitgauss<-: getter and setter for the fitgauss slot of the object.

noise,noise<-: getter and setter for the noise slot of the object.

verboseColumns,verboseColumns<-: getter and setter for the verboseColumns slot of the object.

roiList,roiList<-: getter and setter for the roiList slot of the object.

fistBaselineCheck,firstBaselineCheck<-: getter and setter for the firstBaselineCheck slot
of the object.</pre>

roiScales,roiScales<-: getter and setter for the roiScales slot of the object.

Usage

CentWaveParam(

```
ppm = 25,
  peakwidth = c(20, 50),
  snthresh = 10,
  prefilter = c(3, 100),
 mzCenterFun = "wMean",
  integrate = 1L,
 mzdiff = -0.001,
  fitgauss = FALSE,
  noise = 0,
  verboseColumns = FALSE,
  roiList = list(),
  firstBaselineCheck = TRUE,
 roiScales = numeric(),
 extendLengthMSW = FALSE
)
## S4 method for signature 'OnDiskMSnExp,CentWaveParam'
findChromPeaks(
 object,
 param,
 BPPARAM = bpparam(),
  return.type = "XCMSnExp",
 msLevel = 1L
)
## S4 method for signature 'CentWaveParam'
ppm(object)
## S4 replacement method for signature 'CentWaveParam'
ppm(object) <- value</pre>
## S4 method for signature 'CentWaveParam'
peakwidth(object)
## S4 replacement method for signature 'CentWaveParam'
peakwidth(object) <- value</pre>
## S4 method for signature 'CentWaveParam'
snthresh(object)
## S4 replacement method for signature 'CentWaveParam'
snthresh(object) <- value</pre>
## S4 method for signature 'CentWaveParam'
prefilter(object)
```

findChromPeaks-centWave

S4 replacement method for signature 'CentWaveParam' prefilter(object) <- value</pre> ## S4 method for signature 'CentWaveParam' mzCenterFun(object) ## S4 replacement method for signature 'CentWaveParam' mzCenterFun(object) <- value</pre> ## S4 method for signature 'CentWaveParam' integrate(f) ## S4 replacement method for signature 'CentWaveParam' integrate(object) <- value</pre> ## S4 method for signature 'CentWaveParam' mzdiff(object) ## S4 replacement method for signature 'CentWaveParam' mzdiff(object) <- value</pre> ## S4 method for signature 'CentWaveParam' fitgauss(object) ## S4 replacement method for signature 'CentWaveParam' fitgauss(object) <- value</pre> ## S4 method for signature 'CentWaveParam' noise(object) ## S4 replacement method for signature 'CentWaveParam' noise(object) <- value</pre> ## S4 method for signature 'CentWaveParam' verboseColumns(object) ## S4 replacement method for signature 'CentWaveParam' verboseColumns(object) <- value</pre> ## S4 method for signature 'CentWaveParam' roiList(object) ## S4 replacement method for signature 'CentWaveParam' roiList(object) <- value</pre> ## S4 method for signature 'CentWaveParam' firstBaselineCheck(object)

```
## S4 replacement method for signature 'CentWaveParam'
firstBaselineCheck(object) <- value
## S4 method for signature 'CentWaveParam'
roiScales(object)
## S4 replacement method for signature 'CentWaveParam'
roiScales(object) <- value</pre>
```

Arguments

ppm	numeric(1) defining the maximal tolerated m/z deviation in consecutive scans in parts per million (ppm) for the initial ROI definition.
peakwidth	numeric(2) with the expected approximate peak width in chromatographic space. Given as a range (min, max) in seconds.
snthresh	numeric(1) defining the signal to noise ratio cutoff.
prefilter	numeric(2): $c(k, I)$ specifying the prefilter step for the first analysis step (ROI detection). Mass traces are only retained if they contain at least k peaks with intensity >= I.
mzCenterFun	Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.
integrate	Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.
mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.
noise	numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).
verboseColumns	logical(1) whether additional peak meta data columns should be returned.
roiList	An optional list of regions-of-interest (ROI) representing detected mass traces. If ROIs are submitted the first analysis step is omitted and chromatographic peak detection is performed on the submitted ROIs. Each ROI is expected to have the following elements specified: scmin (start scan index), scmax (end scan index), mzmin (minimum m/z), mzmax (maximum m/z), length (number of scans), intensity (summed intensity). Each ROI should be represented by a list of elements or a single row data.frame.

firstBaselineCheck	
	logical(1). If TRUE continuous data within regions of interest is checked to be above the first baseline.
roiScales	Optional numeric vector with length equal to roiList defining the scale for each region of interest in roiList that should be used for the centWave-wavelets.
extendLengthMSW	l
	Option to force centWave to use all scales when running centWave rather than truncating with the EIC length. Uses the "open" method to extend the EIC to a integer base-2 length prior to being passed to convolve rather than the default "reflect" method. See https://github.com/sneumann/xcms/issues/445 for more information.
object	For findChromPeaks: an ${\tt OnDiskMSnExp}$ object containing the MS- and all other experiment-relevant data.
	For all other methods: a parameter object.
param	An CentWaveParam object containing all settings for the centWave algorithm.
BPPARAM	A parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.
return.type	Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".
msLevel	integer(1) defining the MS level on which the peak detection should be performed. Defaults to $msLevel = 1$.
value	The value for the slot.
f	For integrate: a CentWaveParam object.

Details

The centWave algorithm is most suitable for high resolution LC/{TOF,OrbiTrap,FTICR}-MS data in centroid mode. In the first phase the method identifies *regions of interest* (ROIs) representing mass traces that are characterized as regions with less than ppm m/z deviation in consecutive scans in the LC/MS map. In detail, starting with a single m/z, a ROI is extended if a m/z can be found in the next scan (spectrum) for which the difference to the mean m/z of the ROI is smaller than the user defined ppm of the m/z. The mean m/z of the ROI is then updated considering also the newly included m/z value.

These ROIs are then, after some cleanup, analyzed using continuous wavelet transform (CWT) to locate chromatographic peaks on different scales. The first analysis step is skipped, if regions of interest are passed *via* the param parameter.

Parallel processing (one process per sample) is supported and can be configured either by the BPPARAM parameter or by globally defining the parallel processing mode using the register method from the BiocParallel package.

Value

The CentWaveParam function returns a CentWaveParam class instance with all of the settings specified for chromatographic peak detection by the centWave method.

findChromPeaks-centWave

For findChromPeaks: if return.type = "XCMSnExp" an XCMSnExp object with the results of the peak detection. If return.type = "list" a list of length equal to the number of samples with matrices specifying the identified peaks. If return.type = "xcmsSet" an xcmsSet object with the results of the peak detection.

Slots

ppm,peakwidth,snthresh,prefilter,mzCenterFun,integrate,mzdiff,fitgauss,noise,verboseColumns,roiList See corresponding parameter above. Slots values should exclusively be accessed via the corresponding getter and setter methods listed above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the findPeaks methods. It supports peak detection on OnDiskMSnExp objects (defined in the MSnbase package). All of the settings to the centWave algorithm can be passed with a CentWaveParam object.

Author(s)

Ralf Tautenhahn, Johannes Rainer

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

The do_findChromPeaks_centWave core API function and findPeaks.centWave for the old user interface.

peaksWithCentWave for functions to perform centWave peak detection in purely chromatographic data.

XCMSnExp for the object containing the results of the peak detection.

Other peak detection methods: chromatographic-peak-detection, findChromPeaks-centWaveWithPredIsoROIs, findChromPeaks-massifquant, findChromPeaks-matchedFilter, findPeaks-MSW

Examples

package

```
## Create a CentWaveParam object. Note that the noise is set to 10000 to
## speed up the execution of the example - in a real use case the default
## value should be used, or it should be set to a reasonable value.
cwp <- CentWaveParam(ppm = 20, noise = 10000, prefilter = c(3, 10000))
## Change snthresh parameter
snthresh(cwp) <- 25
cwp
## Perform the peak detection using centWave on some of the files from the
## faahKO package. Files are read using the readMSData from the MSnbase
```

findChromPeaks-centWaveWithPredIsoROIs

Description

This method performs a two-step centWave-based chromatographic peak detection: in a first cent-Wave run peaks are identified for which then the location of their potential isotopes in the mzretention time is predicted. A second centWave run is then performed on these *regions of interest* (ROIs). The final list of chromatographic peaks comprises all non-overlapping peaks from both centWave runs.

The CentWavePredIsoParam class allows to specify all settings for the two-step centWave-based peak detection considering also predicted isotopes of peaks identified in the first centWave run. Instances should be created with the CentWavePredIsoParam constructor. See also the documentation of the CentWaveParam for all methods and arguments this class inherits.

The findChromPeaks, OnDiskMSnExp, CentWavePredIsoParam method performs a two-step centWavebased chromatographic peak detection on all samples from an OnDiskMSnExp object. OnDiskMSnExp objects encapsule all experiment specific data and load the spectra data (mz and intensity values) on the fly from the original files applying also all eventual data manipulations.

snthreshIsoROIs, snthreshIsoROIs<-: getter and setter for the snthreshIsoROIs slot of the object.

maxCharge,maxCharge<-: getter and setter for the maxCharge slot of the object.

maxIso,maxIso<-: getter and setter for the maxIso slot of the object.</pre>

mzIntervalExtension,mzIntervalExtension<-: getter and setter for the mzIntervalExtension
slot of the object.</pre>

polarity,polarity<-: getter and setter for the polarity slot of the object.

Usage

```
CentWavePredIsoParam(
   ppm = 25,
   peakwidth = c(20, 50),
   snthresh = 10,
   prefilter = c(3, 100),
   mzCenterFun = "wMean",
```

```
integrate = 1L,
  mzdiff = -0.001,
  fitgauss = FALSE,
  noise = 0,
  verboseColumns = FALSE,
  roiList = list(),
  firstBaselineCheck = TRUE,
  roiScales = numeric(),
  snthreshIsoROIs = 6.25,
 maxCharge = 3,
 maxIso = 5,
  mzIntervalExtension = TRUE,
  polarity = "unknown"
)
## S4 method for signature 'OnDiskMSnExp,CentWavePredIsoParam'
findChromPeaks(
  object,
  param,
 BPPARAM = bpparam(),
  return.type = "XCMSnExp",
 msLevel = 1L
)
## S4 method for signature 'CentWavePredIsoParam'
snthreshIsoROIs(object)
## S4 replacement method for signature 'CentWavePredIsoParam'
snthreshIsoROIs(object) <- value</pre>
## S4 method for signature 'CentWavePredIsoParam'
maxCharge(object)
## S4 replacement method for signature 'CentWavePredIsoParam'
maxCharge(object) <- value</pre>
## S4 method for signature 'CentWavePredIsoParam'
maxIso(object)
## S4 replacement method for signature 'CentWavePredIsoParam'
maxIso(object) <- value</pre>
## S4 method for signature 'CentWavePredIsoParam'
mzIntervalExtension(object)
## S4 replacement method for signature 'CentWavePredIsoParam'
mzIntervalExtension(object) <- value</pre>
```

S4 method for signature 'CentWavePredIsoParam'
polarity(object)

above the first baseline.

```
## S4 replacement method for signature 'CentWavePredIsoParam'
polarity(object) <- value</pre>
```

Arguments

ppm	numeric(1) defining the maximal tolerated m/z deviation in consecutive scans in parts per million (ppm) for the initial ROI definition.
peakwidth	numeric(2) with the expected approximate peak width in chromatographic space. Given as a range (min, max) in seconds.
snthresh	numeric(1) defining the signal to noise ratio cutoff.
prefilter	numeric(2): $c(k, I)$ specifying the prefilter step for the first analysis step (ROI detection). Mass traces are only retained if they contain at least k peaks with intensity >= I.
mzCenterFun	Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.
integrate	Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.
mzdiff	numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.
noise	numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).
verboseColumns	logical(1) whether additional peak meta data columns should be returned.
roiList	An optional list of regions-of-interest (ROI) representing detected mass traces. If ROIs are submitted the first analysis step is omitted and chromatographic peak detection is performed on the submitted ROIs. Each ROI is expected to have the following elements specified: scmin (start scan index), scmax (end scan index), mzmin (minimum m/z), mzmax (maximum m/z), length (number of scans), intensity (summed intensity). Each ROI should be represented by a list of elements or a single row data.frame.
firstBaselineCheck	
	logical(1). If TRUE continuous data within regions of interest is checked to be

snthreshIsoROISnumeric(1) defining the signal to noise ratio cutoff to be used in the second centWave run to identify peaks for predicted isotope ROIs.maxChargeinteger(1) defining the maximal isotope charge. Isotopes will be defined for charges 1:maxCharge.maxIsointeger(1) defining the number of isotope peaks that should be predicted for each peak identified in the first centWave run.mzIntervalExt=logical(1) whether the mz range for the predicted isotope ROIs should be extended to increase detection of low intensity peaks.polaritycharacter(1) specifying the polarity of the data. Currently not used, but has to be "positive", "negative" or "unknown" if provided.objectFor findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data. For all other methods: a parameter object.BPPARAMA parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocPara1lel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmSSet".msLevelinteger(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.valueThe value for the slot.	roiScales	Optional numeric vector with length equal to roiList defining the scale for each region of interest in roiList that should be used for the centWave-wavelets.
centWave run to identify peaks for predicted isotope ROIs.maxChargeinteger(1) defining the maximal isotope charge. Isotopes will be defined for charges 1:maxCharge.maxIsointeger(1) defining the number of isotope peaks that should be predicted for each peak identified in the first centWave run.mzIntervalExtensionlogical(1) whether the mz range for the predicted isotope ROIs should be extended to increase detection of low intensity peaks.polaritycharacter(1) specifying the polarity of the data. Currently not used, but has to be "positive", "negative" or "unknown" if provided.objectFor findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data. For all other methods: a parameter object.BPPARAMA parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmSSet".msLevelinteger(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.	snthreshIsoROI	S
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each peak identified in the first centWave run.mzIntervalExtensionlogical(1) whether the mz range for the predicted isotope ROIs should be extended to increase detection of low intensity peaks.polaritycharacter(1) specifying the polarity of the data. Currently not used, but has to be "positive", "negative" or "unknown" if provided.objectFor findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data. For all other methods: a parameter object.paramAn CentWavePredIsoParam object with the settings for the chromatographic peak detection algorithm.BPPARAMA parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".msLevelinteger(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.	maxCharge	
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experiment-relevant data.For all other methods: a parameter object.paramAn CentWavePredIsoParam object with the settings for the chromatographic peak detection algorithm.BPPARAMA parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".msLevelinteger(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.	polarity	
paramAn CentWavePredIsoParam object with the settings for the chromatographic peak detection algorithm.BPPARAMA parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".msLevelinteger(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.	object	
peak detection algorithm.BPPARAMA parameter class specifying if and how parallel processing should be performed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.return.typeCharacter specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".msLevelinteger(1) defining the MS level on which the peak detection should be performed. Defaults to msLevel = 1.		For all other methods: a parameter object.
<pre>formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples. return.type Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet". msLevel integer(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.</pre>	param	
"XCMSnExp" (default), "list" or "xcmsSet". msLevel integer(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.	BPPARAM	formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in
formed. Defaults to msLevel = 1.	return.type	
value The value for the slot.	msLevel	
	value	The value for the slot.

Details

See centWave for details on the centWave method.

Parallel processing (one process per sample) is supported and can be configured either by the BPPARAM parameter or by globally defining the parallel processing mode using the register method from the BiocParallel package.

Value

The CentWavePredIsoParam function returns a CentWavePredIsoParam class instance with all of the settings specified for the two-step centWave-based peak detection considering also isotopes.

For findChromPeaks: if return.type = "XCMSnExp" an XCMSnExp object with the results of the peak detection. If return.type = "list" a list of length equal to the number of samples with matrices specifying the identified peaks. If return.type = "xcmsSet" an xcmsSet object with the results of the peak detection.

Slots

ppm,peakwidth,snthresh,prefilter,mzCenterFun,integrate,mzdiff,fitgauss,noise,verboseColumns,roiList See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the findPeaks methods. It supports chromatographic peak detection on OnDiskMSnExp objects (defined in the MSnbase package). All of the settings to the algorithm can be passed with a CentWavePredIsoParam object.

Author(s)

Hendrik Treutler, Johannes Rainer

See Also

The do_findChromPeaks_centWaveWithPredIsoROIs core API function and findPeaks.centWave for the old user interface. CentWaveParam for the class the CentWavePredIsoParam extends.

XCMSnExp for the object containing the results of the peak detection.

Other peak detection methods: chromatographic-peak-detection, findChromPeaks-centWave, findChromPeaks-massifquant, findChromPeaks-matchedFilter, findPeaks-MSW

Examples

```
## Create a param object
p <- CentWavePredIsoParam(maxCharge = 4)
## Change snthresh parameter
snthresh(p) <- 25
p</pre>
```

findChromPeaks-massifquant

Chromatographic peak detection using the massifquant method

Description

Massifquant is a Kalman filter (KF)-based chromatographic peak detection for XC-MS data in centroid mode. The identified peaks can be further refined with the *centWave* method (see findChromPeaks-centWave for details on centWave) by specifying withWave = TRUE.

The MassifquantParam class allows to specify all settings for a chromatographic peak detection using the massifquant method eventually in combination with the centWave algorithm. Instances should be created with the MassifquantParam constructor.

The findChromPeaks, OnDiskMSnExp, MassifquantParam method performs chromatographic peak detection using the *massifquant* algorithm on all samples from an OnDiskMSnExp object. OnDiskMSnExp

objects encapsule all experiment specific data and load the spectra data (mz and intensity values) on the fly from the original files applying also all eventual data manipulations.

ppm,ppm<-: getter and setter for the ppm slot of the object.

peakwidth,peakwidth<-: getter and setter for the peakwidth slot of the object.

snthresh, snthresh<-: getter and setter for the snthresh slot of the object.

prefilter, prefilter <-: getter and setter for the prefilter slot of the object.

mzCenterFun,mzCenterFun<-: getter and setter for the mzCenterFun slot of the object.

integrate, integrate <-: getter and setter for the integrate slot of the object.

mzdiff,mzdiff<-: getter and setter for the mzdiff slot of the object.

fitgauss, fitgauss<-: getter and setter for the fitgauss slot of the object.

noise,noise<-: getter and setter for the noise slot of the object.

verboseColumns,verboseColumns<-: getter and setter for the verboseColumns slot of the object.

criticalValue, criticalValue<-: getter and setter for the criticalValue slot of the object.

consecMissedLimit,consecMissedLimit<-: getter and setter for the consecMissedLimit slot of the object.

unions, unions <-: getter and setter for the unions slot of the object.

checkBack,checkBack<-: getter and setter for the checkBack slot of the object.

withWave,withWave<-: getter and setter for the withWave slot of the object.

Usage

```
MassifquantParam(
  ppm = 25,
  peakwidth = c(20, 50),
  snthresh = 10,
 prefilter = c(3, 100),
 mzCenterFun = "wMean",
  integrate = 1L,
 mzdiff = -0.001,
  fitgauss = FALSE,
  noise = 0,
  verboseColumns = FALSE,
  criticalValue = 1.125,
  consecMissedLimit = 2,
  unions = 1,
  checkBack = 0,
  withWave = FALSE
)
## S4 method for signature 'OnDiskMSnExp,MassifquantParam'
findChromPeaks(
  object,
  param,
 BPPARAM = bpparam(),
```

```
return.type = "XCMSnExp",
 msLevel = 1L
)
## S4 method for signature 'MassifquantParam'
ppm(object)
## S4 replacement method for signature 'MassifquantParam'
ppm(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
peakwidth(object)
## S4 replacement method for signature 'MassifquantParam'
peakwidth(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
snthresh(object)
## S4 replacement method for signature 'MassifquantParam'
snthresh(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
prefilter(object)
## S4 replacement method for signature 'MassifquantParam'
prefilter(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
mzCenterFun(object)
## S4 replacement method for signature 'MassifquantParam'
mzCenterFun(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
integrate(f)
## S4 replacement method for signature 'MassifquantParam'
integrate(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
mzdiff(object)
## S4 replacement method for signature 'MassifquantParam'
mzdiff(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
fitgauss(object)
```

```
## S4 replacement method for signature 'MassifquantParam'
fitgauss(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
noise(object)
## S4 replacement method for signature 'MassifquantParam'
noise(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
verboseColumns(object)
## S4 replacement method for signature 'MassifquantParam'
verboseColumns(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
criticalValue(object)
## S4 replacement method for signature 'MassifquantParam'
criticalValue(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
consecMissedLimit(object)
## S4 replacement method for signature 'MassifquantParam'
consecMissedLimit(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
unions(object)
## S4 replacement method for signature 'MassifquantParam'
unions(object) <- value
## S4 method for signature 'MassifquantParam'
checkBack(object)
## S4 replacement method for signature 'MassifquantParam'
checkBack(object) <- value</pre>
## S4 method for signature 'MassifquantParam'
withWave(object)
## S4 replacement method for signature 'MassifquantParam'
```

withWave(object) <- value</pre>

Arguments

ppm

numeric(1) defining the maximal tolerated m/z deviation in consecutive scans

in parts per	million	(ppm)	for the	initial	ROI	definition.	
in puits per	mminon	(ppm)	ior the	mmuu	nor	deminition.	

peakwidth numeric(2). Only the first element is used by massifquant, which specifices the minimum peak length in time scans. For withWave = TRUE the second argument represents the maximum peak length subject to being greater than the minimum peak length (see also documentation of do_findChromPeaks_centWave).

snthresh numeric(1) defining the signal to noise ratio cutoff.

- prefilter numeric(2). The first argument is only used if (withWave = TRUE); see findChromPeaks-centWave for details. The second argument specifies the minimum threshold for the maximum intensity of a chromatographic peak that must be met.
- mzCenterFun Name of the function to calculate the m/z center of the chromatographic peak. Allowed are: "wMean": intensity weighted mean of the peak's m/z values, "mean": mean of the peak's m/z values, "apex": use the m/z value at the peak apex, "wMeanApex3": intensity weighted mean of the m/z value at the peak apex and the m/z values left and right of it and "meanApex3": mean of the m/z value of the peak apex and the m/z values left and right of it.
- integrate Integration method. For integrate = 1 peak limits are found through descent on the mexican hat filtered data, for integrate = 2 the descent is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.
- mzdiff numeric(1) representing the minimum difference in m/z dimension required for peaks with overlapping retention times; can be negative to allow overlap. During peak post-processing, peaks defined to be overlapping are reduced to the one peak with the largest signal.
- fitgauss logical(1) whether or not a Gaussian should be fitted to each peak. This affects mostly the retention time position of the peak.
- noise numeric(1) allowing to set a minimum intensity required for centroids to be considered in the first analysis step (centroids with intensity < noise are omitted from ROI detection).
- verboseColumns logical(1) whether additional peak meta data columns should be returned.
- criticalValue numeric(1). 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 peak 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 peak in the XC-MS sample have a small mass deviance in ppm error, a smaller critical value might be better and vice versa.

consecMissedLimit

integer (1) Suggested values: (1, 2, 3). While a peak is in the process 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 peak.

unionsinteger (1) set to 1 if apply t-test union on segmentation; set to 0 if no t-test to
be applied on chromatographically continous peaks sharing same m/z range. Ex-
planation: With very few data points, sometimes a Kalman Filter stops tracking
a peak prematurely. Another Kalman Filter is instantiated and begins following

	the rest of the signal. Because tracking is done backwards to forwards, this algo- rithmic defect leaves a real peak divided into two segments or more. With this option turned on, the program identifies segmented peaks and combines them (merges them) into one with a two sample t-test. The potential danger of this option is that some truly distinct peaks may be merged.
checkBack	integer(1) set to 1 if turned on; set to 0 if turned off. The convergence of a Kalman Filter to a peak's precise m/z mapping is very fast, but sometimes it incorporates erroneous centroids as part of a peak (especially early on). The scanBack 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 peak 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.
withWave	logical(1) if TRUE, the peaks identified first with Massifquant are subsequently filtered with the second step of the centWave algorithm, which includes wavelet estimation.
object	For findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data.
	For all other methods: a parameter object.
param	An MassifquantParam object containing all settings for the massifquant algorithm.
BPPARAM	A parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.
return.type	Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".
msLevel	integer(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.
value	The value for the slot.
f	For integrate: a MassifquantParam object.

Details

This algorithm's performance has been tested rigorously on high resolution LC/OrbiTrap, TOF-MS data in centroid mode. Simultaneous kalman filters identify chromatographic peaks 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 peak 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.

Parallel processing (one process per sample) is supported and can be configured either by the BPPARAM parameter or by globally defining the parallel processing mode using the register method from the BiocParallel package.

Value

The MassifquantParam function returns a MassifquantParam class instance with all of the settings specified for chromatographic peak detection by the *massifquant* method.

For findChromPeaks: if return.type = "XCMSnExp" an XCMSnExp object with the results of the peak detection. If return.type = "list" a list of length equal to the number of samples with matrices specifying the identified peaks. If return.type = "xcmsSet" an xcmsSet object with the results of the peak detection.

Slots

ppm, peakwidth, snthresh, prefilter, mzCenterFun, integrate, mzdiff, fitgauss, noise, verboseColumns, critica See corresponding parameter above. Slots values should exclusively be accessed *via* the corresponding getter and setter methods listed above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the findPeaks methods. It supports chromatographic peak detection on OnDiskMSnExp objects (defined in the MSnbase package). All of the settings to the massifquant and centWave algorithm can be passed with a MassifquantParam object.

Author(s)

Christopher Conley, Johannes Rainer

References

Conley CJ, Smith R, Torgrip RJ, Taylor RM, Tautenhahn R and Prince JT "Massifquant: opensource Kalman filter-based XC-MS isotope trace feature detection" *Bioinformatics* 2014, 30(18):2636-43.

See Also

The do_findChromPeaks_massifquant core API function and findPeaks.massifquant for the old user interface.

XCMSnExp for the object containing the results of the peak detection.

Other peak detection methods: chromatographic-peak-detection, findChromPeaks-centWaveWithPredIsoROIs, findChromPeaks-centWave, findChromPeaks-matchedFilter, findPeaks-MSW

Examples

```
## Create a MassifquantParam object.
mqp <- MassifquantParam()
## Change snthresh prefilter parameters
snthresh(mqp) <- 30
prefilter(mqp) <- c(6, 10000)
mqp
```

Perform the peak detection using massifquant on the files from the

findChromPeaks-matchedFilter

Peak detection in the chromatographic time domain

Description

The *matchedFilter* algorithm identifies peaks in the chromatographic time domain as described in [Smith 2006]. The intensity values are binned by cutting The LC/MS data into slices (bins) of a mass unit (binSize m/z) wide. Within each bin the maximal intensity is selected. The chromatographic peak detection is then performed in each bin by extending it based on the steps parameter to generate slices comprising bins current_bin -steps +1 to current_bin + steps -1. Each of these slices is then filtered with matched filtration using a second-derative Gaussian as the model peak shape. After filtration peaks are detected using a signal-to-ratio cut-off. For more details and illustrations see [Smith 2006].

The MatchedFilterParam class allows to specify all settings for a chromatographic peak detection using the matchedFilter method. Instances should be created with the MatchedFilterParam constructor.

The findChromPeaks, OnDiskMSnExp, MatchedFilterParam method performs peak detection using the *matchedFilter* algorithm on all samples from an OnDiskMSnExp object. OnDiskMSnExp objects encapsule all experiment specific data and load the spectra data (mz and intensity values) on the fly from the original files applying also all eventual data manipulations.

binSize,binSize<-: getter and setter for the binSize slot of the object.

impute, impute <-: getter and setter for the impute slot of the object.

baseValue,baseValue<-: getter and setter for the baseValue slot of the object.

distance, distance <-: getter and setter for the distance slot of the object.

fwhm,fwhm<-: getter and setter for the fwhm slot of the object.

sigma, sigma <-: getter and setter for the sigma slot of the object.

max,max<-: getter and setter for the max slot of the object.

snthresh, snthresh <-: getter and setter for the snthresh slot of the object.

steps, steps <-: getter and setter for the steps slot of the object.

mzdiff,mzdiff<-: getter and setter for the mzdiff slot of the object.

index,index<-: getter and setter for the index slot of the object.

Usage

```
MatchedFilterParam(
  binSize = 0.1,
  impute = "none",
  baseValue = numeric(),
  distance = numeric(),
  fwhm = 30,
  sigma = fwhm/2.3548,
  max = 5,
  snthresh = 10,
  steps = 2,
 mzdiff = 0.8 - binSize * steps,
  index = FALSE
)
## S4 method for signature 'OnDiskMSnExp,MatchedFilterParam'
findChromPeaks(
 object,
  param,
 BPPARAM = bpparam(),
  return.type = "XCMSnExp",
 msLevel = 1L
)
## S4 method for signature 'MatchedFilterParam'
binSize(object)
## S4 replacement method for signature 'MatchedFilterParam'
binSize(object) <- value</pre>
## S4 method for signature 'MatchedFilterParam'
impute(object)
## S4 replacement method for signature 'MatchedFilterParam'
impute(object) <- value</pre>
## S4 method for signature 'MatchedFilterParam'
baseValue(object)
## S4 replacement method for signature 'MatchedFilterParam'
baseValue(object) <- value</pre>
## S4 method for signature 'MatchedFilterParam'
distance(object)
## S4 replacement method for signature 'MatchedFilterParam'
distance(object) <- value
```

S4 method for signature 'MatchedFilterParam' fwhm(object) ## S4 replacement method for signature 'MatchedFilterParam' fwhm(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' sigma(object) ## S4 replacement method for signature 'MatchedFilterParam' sigma(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' max(x)## S4 replacement method for signature 'MatchedFilterParam' max(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' snthresh(object) ## S4 replacement method for signature 'MatchedFilterParam' snthresh(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' steps(object) ## S4 replacement method for signature 'MatchedFilterParam' steps(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' mzdiff(object) ## S4 replacement method for signature 'MatchedFilterParam' mzdiff(object) <- value</pre> ## S4 method for signature 'MatchedFilterParam' index(object) ## S4 replacement method for signature 'MatchedFilterParam' index(object) <- value</pre>

Arguments

binSize numeric(1) specifying the width of the bins/slices in m/z dimension.

impute Character string specifying the method to be used for missing value imputation. Allowed values are "none" (no linear interpolation), "lin" (linear interpolation), "linbase" (linear interpolation within a certain bin-neighborhood) and

"intlin". See imputeLinInterpol for more details.

	incline . See impliced introl por for more domins.
baseValue	The base value to which empty elements should be set. This is only considered for method = "linbase" and corresponds to the profBinLinBase's baselevel argument.
distance	For method = "linbase": number of non-empty neighboring element of an empty element that should be considered for linear interpolation. See details section for more information.
fwhm	numeric(1) specifying the full width at half maximum of matched filtration gaussian model peak. Only used to calculate the actual sigma, see below.
sigma	numeric(1) specifying the standard deviation (width) of the matched filtration model peak.
max	numeric(1) representing the maximum number of peaks that are expected/will be identified per slice.
snthresh	numeric(1) defining the signal to noise cutoff to be used in the chromatographic peak detection step.
steps	numeric(1) defining the number of bins to be merged before filtration (i.e. the number of neighboring bins that will be joined to the slice in which filtration and peak detection will be performed).
mzdiff	numeric(1) defining the minimum difference in m/z for peaks with overlapping retention times
index	logical(1) specifying whether indicies should be returned instead of values for m/z and retention times.
object	For findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data.
	For all other methods: a parameter object.
param	An MatchedFilterParam object containing all settings for the matchedFilter algorithm.
BPPARAM	A parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.
return.type	Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".
msLevel	integer(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.
value	The value for the slot.
х	For max: a MatchedFilterParam object.

Details

The intensities are binned by the provided m/z values within each spectrum (scan). Binning is performed such that the bins are centered around the m/z values (i.e. the first bin includes all m/z values between min(mz) -bin_size/2 and min(mz) + bin_size/2).

For more details on binning and missing value imputation see binYonX and imputeLinInterpol methods.

Parallel processing (one process per sample) is supported and can be configured either by the BPPARAM parameter or by globally defining the parallel processing mode using the register method from the BiocParallel package.

Value

The MatchedFilterParam function returns a MatchedFilterParam class instance with all of the settings specified for chromatographic detection by the *matchedFilter* method.

For findChromPeaks: if return.type = "XCMSnExp" an XCMSnExp object with the results of the peak detection. If return.type = "list" a list of length equal to the number of samples with matrices specifying the identified peaks. If return.type = "xcmsSet" an xcmsSet object with the results of the peak detection.

Slots

binSize, impute, baseValue, distance, fwhm, sigma, max, snthresh, steps, mzdiff, index See corresponding parameter above. Slots values should exclusively be accessed *via* the corresponding getter and setter methods listed above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the findPeaks methods. It supports chromatographic peak detection on OnDiskMSnExp objects (defined in the MSnbase package). All of the settings to the matchedFilter algorithm can be passed with a MatchedFilterParam object.

Author(s)

Colin A Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" *Anal. Chem.* 2006, 78:779-787.

See Also

The do_findChromPeaks_matchedFilter core API function and findPeaks.matchedFilter for the old user interface.

peaksWithMatchedFilter for functions to perform matchedFilter peak detection in purely chromatographic data.

XCMSnExp for the object containing the results of the chromatographic peak detection.

Other peak detection methods: chromatographic-peak-detection, findChromPeaks-centWaveWithPredIsoROIs, findChromPeaks-centWave, findChromPeaks-massifquant, findPeaks-MSW

Examples

```
## Create a MatchedFilterParam object. Note that we use a unnecessarily large
## binSize parameter to reduce the run-time of the example.
mfp <- MatchedFilterParam(binSize = 5)</pre>
## Change snthresh parameter
snthresh(mfp) <- 15
mfp
## Perform the peak detection using matchecFilter on the files from the
## faahKO package. Files are read using the readMSData from the MSnbase
## package
library(faahKO)
library(MSnbase)
fls <- dir(system.file("cdf/KO", package = "faahKO"), recursive = TRUE,</pre>
           full.names = TRUE)
raw_data <- readMSData(fls[1], mode = "onDisk")</pre>
## Perform the chromatographic peak detection using the settings defined
## above. Note that we are also disabling parallel processing in this
## example by registering a "SerialParam"
res <- findChromPeaks(raw_data, param = mfp)</pre>
head(chromPeaks(res))
```

findChromPeaksIsolationWindow

Data independent acquisition (DIA): peak detection in isolation windows

Description

The findChromPeaksIsolationWindow function allows to perform a chromatographic peak detection in MS level > 1 spectra of certain isolation windows (e.g. SWATH pockets). The function performs a peak detection, separately for all spectra belonging to the same isolation window and adds them to the chromPeaks() matrix of the result object, information about the isolation window they were detected in is added to chromPeakData(). Note that peak detection with this method does not remove previously identified chromatographic peaks (e.g. on MS1 level using the findChromPeaks() function but adds newly identified peaks to the existing chromPeaks() matrix.

Isolation windows can be defined with the isolationWindow parameter, that by default uses the definition of isolationWindowTargetMz(), i.e. chromatographic peak detection is performed for all spectra with the same isolation window target m/z (seprarately for each file). The parameter param allows to define and configure the peak detection algorithm (see findChromPeaks() for more information).

Usage

```
findChromPeaksIsolationWindow(
   object,
   param,
   msLevel = 2L,
```

```
isolationWindow = isolationWindowTargetMz(object),
...
```

Arguments

)

object	OnDiskMSnExp or XCMSnExp object with the DIA data.	
param	Peak detection parameter object, such as a CentWaveParam object defining and configuring the chromographic peak detection algorithm. See also findChromPeaks() for more details.	
msLevel	integer(1) specifying the MS level in which the peak detection should be per- formed. By default msLevel = 2L.	
isolationWindow		
	factor or similar defining the isolation windows in which the peak detection should be performed with length equal to the number of spectra in object.	
	currently not used.	

Value

An XCMSnExp object with the chromatographic peaks identified in spectra of each isolation window from each file added to the chromPeaks matrix. Isolation window definition for each identified peak are stored as additional columns in chromPeakData().

Author(s)

Johannes Rainer, Michael Witting

See Also

reconstructChromPeakSpectra() for the function to reconstruct MS2 spectra for each MS1 chromatographic peak.

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)

findMZ

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	D
	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, <hpaul.beonton08@imperial.ac.uk>

References

H. Paul Benton, D.M. Wong, S.A.Strauger, G. Siuzdak " $XCMS^2$ " Analytical Chemistry 2008

See Also

findneutral,

Examples

```
##takes only one file from the file set
xfrag <- xcmsFragments(xs)
found<-findMZ(xfrag, 657.3433, 50)</pre>
```

End(Not run)

findneutral

Find neutral losses in xcmsFragment objects

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
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

findPeaks-methods

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

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.

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 findPeaks.addPredictedIsotopeFeatures
findPeaks.centWaveWithPredictedIsotopeROIs xcmsRaw-class

findPeaks-MSW Single-spectrum non-chromatography MS data peak detection

Description

Perform peak detection in mass spectrometry direct injection spectrum using a wavelet based algorithm.

The MSWParam class allows to specify all settings for a peak detection using the MSW method. Instances should be created with the MSWParam constructor.

The findChromPeaks, OnDiskMSnExp, MSWParam method performs peak detection in single-spectrum non-chromatography MS data using functionality from the MassSpecWavelet package on all samples from an OnDiskMSnExp object. OnDiskMSnExp objects encapsule all experiment specific data and load the spectra data (mz and intensity values) on the fly from the original files applying also all eventual data manipulations.

snthresh, snthresh<-: getter and setter for the snthresh slot of the object.

verboseColumns,verboseColumns<-: getter and setter for the verboseColumns slot of the object.

scales, scales <-: getter and setter for the scales slot of the object.

nearbyPeak,nearbyPeak<-: getter and setter for the nearbyPeak slot of the object.

peakScaleRange,peakScaleRange<-: getter and setter for the peakScaleRange slot of the object.

ampTh,ampTh<-: getter and setter for the ampTh slot of the object.</pre>

minNoiseLevel,minNoiseLevel<-: getter and setter for the minNoiseLevel slot of the object.

ridgeLength,ridgeLength<-: getter and setter for the ridgeLength slot of the object.

peakThr,peakThr<-: getter and setter for the peakThr slot of the object.

tuneIn,tuneIn<-: getter and setter for the tuneIn slot of the object.</pre>

addParams,addParams<-: getter and setter for the addParams slot of the object. This slot stores optional additional parameters to be passed to the identifyMajorPeaks and sav.gol functions from the MassSpecWavelet package.

Usage

```
MSWParam(
  snthresh = 3,
  verboseColumns = FALSE,
  scales = c(1, seq(2, 30, 2), seq(32, 64, 4)),
  nearbyPeak = TRUE,
  peakScaleRange = 5,
  ampTh = 0.01,
 minNoiseLevel = ampTh/snthresh,
  ridgeLength = 24,
  peakThr = NULL,
  tuneIn = FALSE,
  . . .
)
## S4 method for signature 'OnDiskMSnExp,MSWParam'
findChromPeaks(
  object,
  param,
 BPPARAM = bpparam(),
  return.type = "XCMSnExp",
  msLevel = 1L
)
## S4 method for signature 'MSWParam'
snthresh(object)
## S4 replacement method for signature 'MSWParam'
snthresh(object) <- value</pre>
## S4 method for signature 'MSWParam'
verboseColumns(object)
## S4 replacement method for signature 'MSWParam'
verboseColumns(object) <- value</pre>
## S4 method for signature 'MSWParam'
scales(object)
```

```
## S4 replacement method for signature 'MSWParam'
scales(object) <- value</pre>
## S4 method for signature 'MSWParam'
nearbyPeak(object)
## S4 replacement method for signature 'MSWParam'
nearbyPeak(object) <- value</pre>
## S4 method for signature 'MSWParam'
peakScaleRange(object)
## S4 replacement method for signature 'MSWParam'
peakScaleRange(object) <- value</pre>
## S4 method for signature 'MSWParam'
ampTh(object)
## S4 replacement method for signature 'MSWParam'
ampTh(object) <- value</pre>
## S4 method for signature 'MSWParam'
minNoiseLevel(object)
## S4 replacement method for signature 'MSWParam'
minNoiseLevel(object) <- value</pre>
## S4 method for signature 'MSWParam'
ridgeLength(object)
## S4 replacement method for signature 'MSWParam'
ridgeLength(object) <- value</pre>
## S4 method for signature 'MSWParam'
peakThr(object)
## S4 replacement method for signature 'MSWParam'
peakThr(object) <- value</pre>
## S4 method for signature 'MSWParam'
tuneIn(object)
## S4 replacement method for signature 'MSWParam'
tuneIn(object) <- value</pre>
## S4 method for signature 'MSWParam'
addParams(object)
```

S4 replacement method for signature 'MSWParam'
addParams(object) <- value</pre>

Arguments

snthresh	numeric(1) defining the signal to noise ratio cutoff.
verboseColumns	logical(1) whether additional peak meta data columns should be returned.
scales	Numeric defining the scales of the continuous wavelet transform (CWT).
nearbyPeak	logical(1) whether to include nearby peaks of major peaks.
peakScaleRange	numeric(1) defining the scale range of the peak (larger than 5 by default).
ampTh	numeric(1) defining the minimum required relative amplitude of the peak (ratio of the maximum of CWT coefficients).
minNoiseLevel	numeric(1) defining the minimum noise level used in computing the SNR.
ridgeLength	numeric(1) defining the minimum highest scale of the peak in 2-D CWT coefficient matrix.
peakThr	numeric(1) with the minimum absolute intensity (above baseline) of peaks to be picked. If provided, the smoothing function sav.gol function (in the MassSpecWavelet) package is called to estimate the local intensity.
tuneIn	logical(1) when to tune in the parameter estimation of the detected peaks.
	Additional parameters to be passed to the identifyMajorPeaks and sav.gol functions from the MassSpecWavelet package.
object	For findChromPeaks: an OnDiskMSnExp object containing the MS- and all other experiment-relevant data.
	For all other methods: a parameter object.
param	An MSWParam object containing all settings for the algorithm.
BPPARAM	A parameter class specifying if and how parallel processing should be per- formed. It defaults to bpparam. See documentation of the BiocParallel for more details. If parallel processing is enabled, peak detection is performed in parallel on several of the input samples.
return.type	Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".
msLevel	integer(1) defining the MS level on which the peak detection should be per- formed. Defaults to msLevel = 1.
value	The value for the slot.

Details

This is a wrapper for the peak picker in Bioconductor's MassSpecWavelet package calling peakDetectionCWT and tuneInPeakInfo functions. See the *xcmsDirect* vignette for more information.

Parallel processing (one process per sample) is supported and can be configured either by the BPPARAM parameter or by globally defining the parallel processing mode using the register method from the BiocParallel package.

The MSWParam function returns a MSWParam class instance with all of the settings specified for peak detection by the *MSW* method.

For findChromPeaks: if return.type = "XCMSnExp" an XCMSnExp object with the results of the peak detection. If return.type = "list" a list of length equal to the number of samples with matrices specifying the identified peaks. If return.type = "xcmsSet" an xcmsSet object with the results of the detection.

Slots

snthresh,verboseColumns,scales,nearbyPeak,peakScaleRange,ampTh,minNoiseLevel,ridgeLength,peakThr,tu See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the findPeaks methods. It supports peak detection on OnDiskMSnExp objects (defined in the MSnbase package). All of the settings to the algorithm can be passed with a MSWParam object.

Author(s)

Joachim Kutzera, Steffen Neumann, Johannes Rainer

See Also

The do_findPeaks_MSW core API function and findPeaks.MSW for the old user interface.

XCMSnExp for the object containing the results of the peak detection.

Other peak detection methods: chromatographic-peak-detection, findChromPeaks-centWaveWithPredIsoROIs, findChromPeaks-centWave, findChromPeaks-massifquant, findChromPeaks-matchedFilter

Examples

head(chromPeaks(fticr))

```
{\tt find Peaks.add Predicted Isotope Features-methods}
```

Feature detection based on predicted isotope features for high resolution LC/MS data

Description

Peak density and wavelet based feature detection aiming at isotope peaks 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	
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.column	-	
	logical, if TRUE additional peak meta data columns are returned	
xcmsPeaks	peak list picked using the centWave algorithm with parameter verbose.columns set to TRUE (columns scmin and scmax needed)	
snthresh	signal to noise ratio cutoff, definition see below.	
maxcharge	max. number of the isotope charge.	
maxiso	max. number of the isotope peaks to predict for each detected feature.	
mzIntervalExtension		
	logical, if TRUE predicted isotope ROIs (regions of interest) are extended in the m/z dimension to increase the detection of low intensity and hence noisy peaks.	

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 isotope ROIs (regions of interest) in the LC/MS map are predicted. In the second phase these mass traces are further analysed. Continuous wavelet transform (CWT) is used to locate chromatographic peaks on different scales. The resulting peak list and the given peak list (xcmsPeaks) are merged and redundant peaks are removed.

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.
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.centWave(object,ppm=25,peakwidth=c(20,50),prefilter=c(3,100),mzCenterF numeric(),noise=0,sleep=0,verbose.columns=FALSE,xcmsPeaks,snthresh=6.25,maxcharge=3,maxiso=5,mz

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\ Hendrik Treutler and Steffen Neumann. "Prediction, detection, and validation of isotope clusters in mass spectrometry data" Submitted to Metabolites 2016, Special Issue "Bioinformatics and Data Analysis"

See Also

findPeaks.centWave findPeaks-methods 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.	
firstBaselineCheck		
	logical, if TRUE continuous data within ROI is checked to be above 1st baseline	
roiScales	numeric, optional vector of scales for each ROI in $\tt ROI.list$ to be used for the <code>centWave-wavelets</code>	

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.
egauss	RMSE of Gaussian fit
	if verbose.columns is TRUE additionally :
mu	Gaussian parameter mu

findPeaks.centWaveWithPredictedIsotopeROIs-methods

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.centWave(object,ppm=25,peakwidth=c(20,50),snthresh=10,prefilter=c(3,10
 numeric(),noise=0,sleep=0,verbose.columns=FALSE,ROI.list=list()),firstBaselineCheck=TRUE,roiSca

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

centWave for the new user interface. findPeaks-methods xcmsRaw-class

Description

Peak density and wavelet based feature detection for high resolution LC/MS data in centroid mode with additional peak picking of isotope features on basis of isotope peak predictions

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 \geq 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.column	
	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.
firstBaselineC	
	logical, if TRUE continuous data within ROI is checked to be above 1st baseline
roiScales	numeric, optional vector of scales for each ROI in ROI.list to be used for the centWave-wavelets
snthreshIsoROI	
	signal to noise ratio cutoff for predicted isotope ROIs, definition see below.
maxcharge	max. number of the isotope charge.
maxiso mzIntervalExte	max. number of the isotope peaks to predict for each detected feature.
	logical, if TRUE predicted isotope ROIs (regions of interest) are extended in the
	m/z dimension to increase the detection of low intensity and hence noisy peaks.

Details

This algorithm is most suitable for high resolution LC/{TOF,OrbiTrap,FTICR}-MS data in centroid mode. The centWave algorithm is applied in two peak picking steps as follows. In the first peak picking step ROIs (regions of interest, characterised as regions with less than ppm m/z deviation in consecutive scans) in the LC/MS map are located and further analysed using continuous wavelet transform (CWT) for the localization of chromatographic peaks on different scales. In the second peak picking step isotope ROIs in the LC/MS map are predicted further analysed using continuous wavelet transform (CWT) for the localization of chromatographic peaks on different scales. The peak lists resulting from both peak picking steps are merged and redundant peaks are removed.

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.
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.centWaveWithPredictedIsotopeROIs(object,ppm=25,peakwidth=c(20,50),snth numeric(),noise=0,sleep=0,verbose.columns=FALSE,ROI.list=list(),firstBaselineCheck=TRUE,roiScal

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\ Hendrik Treutler and Steffen Neumann. "Prediction, detection, and validation of isotope clusters in mass spectrometry data" Submitted to Metabolites 2016, Special Issue "Bioinformatics and Data Analysis"

See Also

do_findChromPeaks_centWaveWithPredIsoROIs for the corresponding core API function. findPeaks.addPredictedIso
findPeaks.centWave findPeaks-methods xcmsRaw-class

findPeaks.massifquant-methods

Feature detection for XC-MS data.

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.

objectalValue 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.

consecMissedLimit

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 mininum 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 incorpo- rates 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 iden- tification 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,prefilter=c(3
    numeric(),noise=0,sleep=0,verbose.columns=FALSE,criticalValue = 1.125,consecMissedLimit
    = 2,unions = 1,checkBack = 0,withWave = 0)
```

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

centWave for the new user interface. findPeaks-methods xcmsSet xcmsRaw xcmsRaw-class

Examples

```
library(faahKO)
library(xcms)
#load all the wild type and Knock out samples
cdfpath <- system.file("cdf", package = "faahKO")</pre>
## Subset to only the first 2 files.
cdffiles <- list.files(cdfpath, recursive = TRUE, full.names = TRUE)[1:2]
## Run the massifquant analysis. Setting the noise level to 10000 to speed up
## execution of the examples - in a real use case it should be set to a reasoable
## value.
xset <- xcmsSet(cdffiles, method = "massifquant",</pre>
                consecMissedLimit = 1,
                snthresh = 10,
                criticalValue = 1.73,
                ppm = 10,
                peakwidth= c(30, 60),
                prefilter= c(1,3000),
                noise = 10000,
                with Wave = 0)
```

Description

Find peaks in the chromatographic time domain of the profile matrix. For more details see do_findChromPeaks_matchedFil

Usage

```
## S4 method for signature 'xcmsRaw'
findPeaks.matchedFilter(
   object,
   fwhm = 30,
   sigma = fwhm/2.3548,
   max = 5,
   snthresh = 10,
   step = 0.1,
   steps = 2,
   mzdiff = 0.8 - step * steps,
   index = FALSE,
   sleep = 0,
   scanrange = numeric()
)
```

Arguments

object	The xcmsRaw object on which peak detection should be performed.
fwhm	numeric(1) specifying the full width at half maximum of matched filtration gaussian model peak. Only used to calculate the actual sigma, see below.
sigma	numeric(1) specifying the standard deviation (width) of the matched filtration model peak.
max	numeric(1) representing the maximum number of peaks that are expected/will be identified per slice.
snthresh	numeric(1) defining the signal to noise cutoff to be used in the chromatographic peak detection step.
step	numeric(1) specifying the width of the bins/slices in m/z dimension.
steps	numeric(1) defining the number of bins to be merged before filtration (i.e. the number of neighboring bins that will be joined to the slice in which filtration and peak detection will be performed).
mzdiff	<code>numeric(1)</code> defining the minimum difference in m/z for peaks with overlapping retention times
index	logical(1) specifying whether indicies should be returned instead of values for m/z and retention times.
sleep	(DEPRECATED). The use of this parameter is highly discouraged, as it could cause problems in parallel processing mode.
scanrange	Numeric vector defining the range of scans to which the original object should be sub-setted before peak detection.

Value

A matrix, each row representing an intentified chromatographic peak, with columns:

mz Intensity weighted mean of m/z values of the peak across scans.

mzmin Minimum m/z of the peak.

mzmax Maximum m/z of the peak.

rt Retention time of the peak's midpoint.

rtmin Minimum retention time of the peak.

rtmax Maximum retention time of the peak.

into Integrated (original) intensity of the peak.

intf Integrated intensity of the filtered peak.

maxo Maximum intensity of the peak.

maxf Maximum intensity of the filtered peak.

i Rank of peak in merged EIC (<= max).

sn Signal to noise ratio of the peak.

Author(s)

Colin A. Smith

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" *Anal. Chem.* 2006, 78:779-787. @family Old peak detection methods

See Also

matchedFilter for the new user interface. xcmsRaw, do_findChromPeaks_matchedFilter for the core function performing the peak detection.

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 precursorIntensity="125245"
</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

into, 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

findPeaks.MSW,xcmsRaw-method

Peak detection for single-spectrum non-chromatography MS data

Description

This method performs peak detection in mass spectrometry direct injection spectrum using a wavelet based algorithm.

Usage

```
## S4 method for signature 'xcmsRaw'
findPeaks.MSW(object, snthresh = 3, verbose.columns = FALSE, ...)
```

Arguments

object	The xcmsRaw object on which peak detection should be performed.	
snthresh	numeric(1) defining the signal to noise ratio cutoff.	
verbose.columns		
	Logical whether additional peak meta data columns should be returned.	
	Additional parameters to be passed to the identifyMajorPeaks and sav.gol functions from the MassSpecWavelet package.	
	runeuons nom the hassopeendveret package.	

Details

This is a wrapper around the peak picker in Bioconductor's MassSpecWavelet package calling peakDetectionCWT and tuneInPeakInfo functions.

Value

A matrix, each row representing an intentified peak, with columns:

mz m/z value of the peak at the centroid position.

mzmin Minimum m/z of the peak.

mzmax Maximum m/z of the peak.

rt Always -1.

rtmin Always -1.

rtmax Always -1.

into Integrated (original) intensity of the peak.

maxo Maximum intensity of the peak.

intf Always NA.

maxf Maximum MSW-filter response of the peak.

sn Signal to noise ratio.

GenericParam-class

Author(s)

Joachim Kutzera, Steffen Neumann, Johannes Rainer

See Also

MSW for the new user interface, do_findPeaks_MSW for the downstream analysis function or peakDetectionCWT from the MassSpecWavelet for details on the algorithm and additionally supported parameters.

GenericParam-class Generic parameter class

Description

The GenericParam class allows to store generic parameter information such as the name of the function that was/has to be called (slot fun) and its arguments (slot args). This object is used to track the process history of the data processings of an XCMSnExp object. This is in contrast to e.g. the CentWaveParam object that is passed to the actual processing method.

Usage

```
GenericParam(fun = character(), args = list())
```

Arguments

fun	character representing the name of the function.
args	list (ideally named) with the arguments to the function.

Value

The GenericParam function returns a GenericParam object.

Slots

fun character specifying the function name.
args list (ideally named) with the arguments to the function.

Author(s)

Johannes Rainer

See Also

processHistory for how to access the process history of an XCMSnExp object.

Examples

```
prm <- GenericParam(fun = "mean")</pre>
```

prm <- GenericParam(fun = "mean", args = list(na.rm = TRUE))</pre>

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.

Note that this method will *always* return profile, not raw data (with profile data being the binned data along M/Z). See details for further information.

Arguments

object	the xcmsRaw or xcmsSet object
mzrange	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 the group data will be used if mzrange is not provided.
rtrange	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	step (bin) size to use for profile generation. Note that a value of $step = 0$ is not supported.
groupidx	either character vector with names or integer vector with indicies of peak groups for which to get EICs
sampleidx	either character vector with names or integer vector with indicies of samples for which to get EICs
rt	"corrected" for using corrected retention times, or "raw" for using raw retention times

Details

In contrast to the rawEIC method, that extracts the actual raw values, this method extracts them from the object's profile matrix (or if the provided step argument does not match the profStep of the object the profile matrix is calculated on the fly and the values returned).

Value

For xcmsSet and xcmsRaw objects, an xcmsEIC object.

getPeaks-methods

Methods

See Also

xcmsRaw-class, xcmsSet-class, xcmsEIC-class, rawEIC

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

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=profinfo
)
```

Author(s)

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

See Also

xcmsRaw-class,

group-methods Grou

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,

150

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,sleep = 0)

See Also

do_groupChromPeaks_density for the core API function performing the analysis. 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)
    xset<-xcmsSet(cdffiles)
    gxset<-group(xset, method="nearest")
    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))),</pre>
                             function(x, object, mf){
                                 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(object)) ))
                                 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]</pre>
    gxset.post@groups<-gxset@groups[idx,]</pre>
    nrow(gxset.post@groups) == 465 ## number of features after minFrac
## End(Not run)
```

groupChromPeaks Correspondence: Chromatographic peak grouping methods.

Description

The groupChromPeaks method(s) perform the correspondence, i.e. the grouping of chromatographic peaks within and between samples. These methods are part of the modernized xcms user interface. The resulting peak groups are referred to as (mz-rt) features and can be accessed *via* the featureDefinitions method on the result object.

The implemented peak grouping methods are:

- **density** peak grouping based on time dimension peak densities. See groupChromPeaks-density for more details.
- **mzClust** high resolution peak grouping for single spectra (direct infusion) MS data. See groupChromPeaks-mzClust for more details.
- **nearest** chromatographic peak grouping based on their proximity in the mz-rt space. See groupChromPeaks-nearest for more details.

Author(s)

Johannes Rainer

See Also

featureDefinitions and featureValues, XCMSnExp-method for methods to access peak grouping results.

featureChromatograms to extract ion chromatograms for each feature.

group for the *old* peak grouping methods.

Other peak grouping methods: groupChromPeaks-density, groupChromPeaks-mzClust, groupChromPeaks-nearest

groupChromPeaks-density

Peak grouping based on time dimension peak densities

Description

This method performs performs correspondence (chromatographic peak grouping) based on the density (distribution) of identified peaks along the retention time axis within slices of overlapping mz ranges. All peaks (from the same or from different samples) being close on the retention time axis are grouped into a feature (*peak group*).

The PeakDensityParam class allows to specify all settings for the peak grouping based on peak densities along the time dimension. Instances should be created with the PeakDensityParam() constructor.

sampleGroups, sampleGroups<-: getter and setter for the sampleGroups slot of the object. Its length should match the number of samples in the experiment and it should not contain NAs.

bw,bw<-: getter and setter for the bw slot of the object.

minFraction,minFraction<-: getter and setter for the minFraction slot of the object.

minSamples, minSamples <-: getter and setter for the minSamples slot of the object.

binSize, binSize <-: getter and setter for the binSize slot of the object.

maxFeatures,maxFeatures<-: getter and setter for the maxFeatures slot of the object. groupChromPeaks,XCMSnExp,PeakDensityParam: performs correspondence (peak grouping within and across samples) within in mz dimension overlapping slices of MS data based on the density distribution of the identified chromatographic peaks in the slice along the time axis.

The correspondence analysis can be performed on chromatographic peaks of any MS level (if present and if chromatographic peak detection has been performed for that MS level) defining features combining these peaks. The MS level can be selected with the parameter msLevel. By default, calling groupChromPeaks will remove any previous correspondence results. This can be disabled with add = TRUE, which will add newly defined features to already present feature definitions.

Usage

```
PeakDensityParam(
  sampleGroups = numeric(),
  bw = 30,
  minFraction = 0.5,
  minSamples = 1,
```

groupChromPeaks-density

```
binSize = 0.25,
  maxFeatures = 50
)
## S4 method for signature 'PeakDensityParam'
sampleGroups(object)
## S4 replacement method for signature 'PeakDensityParam'
sampleGroups(object) <- value</pre>
## S4 method for signature 'PeakDensityParam'
bw(object)
## S4 replacement method for signature 'PeakDensityParam'
bw(object) <- value</pre>
## S4 method for signature 'PeakDensityParam'
minFraction(object)
## S4 replacement method for signature 'PeakDensityParam'
minFraction(object) <- value</pre>
## S4 method for signature 'PeakDensityParam'
minSamples(object)
## S4 replacement method for signature 'PeakDensityParam'
minSamples(object) <- value</pre>
## S4 method for signature 'PeakDensityParam'
binSize(object)
## S4 replacement method for signature 'PeakDensityParam'
binSize(object) <- value</pre>
## S4 method for signature 'PeakDensityParam'
maxFeatures(object)
## S4 replacement method for signature 'PeakDensityParam'
maxFeatures(object) <- value</pre>
## S4 method for signature 'XCMSnExp,PeakDensityParam'
```

groupChromPeaks(object, param, msLevel = 1L, add = FALSE)

Arguments

sampleGroups A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the

156

	same group).
bw	numeric(1) defining the bandwidth (standard deviation of the smoothing ker- nel) to be used. This argument is passed to the [density() method.
minFraction	numeric(1) defining the minimum fraction of samples in at least one sample group in which the peaks have to be present to be considered as a peak group (feature).
minSamples	numeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).
binSize	numeric(1) defining the size of the overlapping slices in mz dimension.
maxFeatures	numeric(1) with the maximum number of peak groups to be identified in a single mz slice.
object	For groupChromPeaks: an XCMSnExp object containing the results from a pre- vious peak detection analysis (see findChromPeaks()).
	For all other methods: a `PeakDensityParam` object.
value	The value for the slot.
param	A PeakDensityParam object containing all settings for the peak grouping algorithm.
msLevel	integer(1) (default msLevel = 1L) defining the MS level on which the corre- spondence should be performed. It is required that chromatographic peaks of the respective MS level are present.
add	logical(1) (default add = FALSE) allowing to perform an additional round of correspondence (e.g. on a different MS level) and add features to the already present feature definitions.

Value

The PeakDensityParam function returns a PeakDensityParam class instance with all of the settings specified for chromatographic peak alignment based on peak densities. Note that argument sampleGroups is mandatory and should represent either the sample grouping in the experiment. It's length has to match the number of sample in the experiments.

For groupChromPeaks: a XCMSnExp object with the results of the correspondence analysis. The definition of the resulting mz-rt features can be accessed with the featureDefinitions() method

Slots

sampleGroups,bw,minFraction,minSamples,binSize,maxFeatures See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface. All of the settings to the algorithm can be passed with a PeakDensityParam object.

Author(s)

Colin Smith, Johannes Rainer

References

Colin A. Smith, Elizabeth J. Want, Grace O'Maille, Ruben Abagyan and Gary Siuzdak. "XCMS: Processing Mass Spectrometry Data for Metabolite Profiling Using Nonlinear Peak Alignment, Matching, and Identification" Anal. Chem. 2006, 78:779-787.

See Also

The do_groupChromPeaks_density() core API function and group.density() for the old user interface.

plotChromPeakDensity() to plot peak densities and evaluate different algorithm settings.

featureDefinitions() and featureValues() for methods to access the features (i.e. the peak grouping results).

XCMSnExp for the object containing the results of the correspondence.

plotChromPeakDensity() for plotting chromatographic peak density with the possibility to test different parameter settings.

Other peak grouping methods: groupChromPeaks-mzClust, groupChromPeaks-nearest, groupChromPeaks()

Examples

```
## Create a PeakDensityParam object
p <- PeakDensityParam(binSize = 0.05, sampleGroups = c(1, 1, 2, 2))</pre>
## Change hte minSamples slot
minSamples(p) <- 3
р
## Chromatographic peak detection and grouping.
##
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")</pre>
## Disable parallel processing for this example
register(SerialParam())
res <- faahko_sub
head(chromPeaks(res))
## The number of peaks identified per sample:
table(chromPeaks(res)[, "sample"])
## Performing the chromatographic peak grouping. Assigning all samples to
## the same sample group.
fdp <- PeakDensityParam(sampleGroups = rep(1, length(fileNames(res))))</pre>
res <- groupChromPeaks(res, fdp)</pre>
## The definition of the features (peak groups):
featureDefinitions(res)
```

```
## Using the featureValues method to extract a matrix with the
## intensities of the features per sample.
head(featureValues(res, value = "into"))
```

```
## The process history:
processHistory(res)
```

groupChromPeaks-mzClust

High resolution peak grouping for single spectra samples

Description

This method performs high resolution correspondence for single spectra samples.

The MzClustParam class allows to specify all settings for the peak grouping based on the *mzClust* algorithm. Instances should be created with the MzClustParam constructor.

sampleGroups,sampleGroups<-: getter and setter for the sampleGroups slot of the object.

ppm,ppm<-: getter and setter for the ppm slot of the object.

absMz,absMz<-: getter and setter for the absMz slot of the object.

minFraction, minFraction <-: getter and setter for the minFraction slot of the object.

minSamples,minSamples<-: getter and setter for the minSamples slot of the object. groupChromPeaks,XCMSnExp,MzClustParam: performs high resolution peak grouping for single spectrum metabolomics data.

Usage

```
MzClustParam(
  sampleGroups = numeric(),
  ppm = 20,
  absMz = 0,
  minFraction = 0.5,
  minSamples = 1
)
## S4 method for signature 'MzClustParam'
sampleGroups(object)
## S4 replacement method for signature 'MzClustParam'
sampleGroups(object) <- value
## S4 method for signature 'MzClustParam'
ppm(object)
## S4 replacement method for signature 'MzClustParam'
```

```
ppm(object) <- value
## S4 method for signature 'MzClustParam'
absMz(object)
## S4 replacement method for signature 'MzClustParam'
absMz(object) <- value
## S4 method for signature 'MzClustParam'
minFraction(object)
## S4 replacement method for signature 'MzClustParam'
minFraction(object) <- value
## S4 method for signature 'MzClustParam'
minSamples(object)
## S4 replacement method for signature 'MzClustParam'
## S4 method for signature 'MzClustParam'</pre>
```

groupChromPeaks(object, param, msLevel = 1L)

```
Arguments
```

ppmnumeric(1) representing the relative mz error for the clustering/grouping (in parts per million).absMznumeric(1) representing the absolute mz error for the clustering.minFractionnumeric(1) defining the minimum fraction of samples in at least one sample group in which the peaks have to be present to be considered as a peak group (feature).minSamplesnumeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).objectFor groupChromPeaks: an XCMSnExp object containing the results from a pre- vious chromatographic peak detection analysis (see findChromPeaks()). For all other methods: a `MzClustParam` object.valueThe value for the slot.paramA MzClustParam object containing all settings for the peak grouping algorithm. integer(1) defining the MS level. Currently only MS level 1 is supported.	sampleGroups	A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the same group).
minFractionnumeric(1) defining the minimum fraction of samples in at least one sample group in which the peaks have to be present to be considered as a peak group (feature).minSamplesnumeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).objectFor groupChromPeaks: an XCMSnExp object containing the results from a pre- vious chromatographic peak detection analysis (see findChromPeaks()). For all other methods: a `MzClustParam` object.valueThe value for the slot.paramA MzClustParam object containing all settings for the peak grouping algorithm.	ppm	
group in which the peaks have to be present to be considered as a peak group (feature).minSamplesnumeric(1) with the minimum number of samples in at least one sample group in which the peaks have to be detected to be considered a peak group (feature).objectFor groupChromPeaks: an XCMSnExp object containing the results from a pre- vious chromatographic peak detection analysis (see findChromPeaks()). For all other methods: a `MzClustParam` object.valueThe value for the slot.paramA MzClustParam object containing all settings for the peak grouping algorithm.	absMz	numeric(1) representing the absolute mz error for the clustering.
 in which the peaks have to be detected to be considered a peak group (feature). bject For groupChromPeaks: an XCMSnExp object containing the results from a previous chromatographic peak detection analysis (see findChromPeaks()). For all other methods: a `MzClustParam` object. value The value for the slot. param A MzClustParam object containing all settings for the peak grouping algorithm. 	minFraction	group in which the peaks have to be present to be considered as a peak group
 vious chromatographic peak detection analysis (see findChromPeaks()). For all other methods: a `MzClustParam` object. value The value for the slot. param A MzClustParam object containing all settings for the peak grouping algorithm. 	minSamples	
valueThe value for the slot.paramA MzClustParam object containing all settings for the peak grouping algorithm.	object	
param A MzClustParam object containing all settings for the peak grouping algorithm.		For all other methods: a `MzClustParam` object.
	value	The value for the slot.
msLevel integer(1) defining the MS level. Currently only MS level 1 is supported.	param	A MzClustParam object containing all settings for the peak grouping algorithm.
	msLevel	integer(1) defining the MS level. Currently only MS level 1 is supported.

160

Value

The MzClustParam function returns a MzClustParam class instance with all of the settings specified for high resolution single spectra peak alignment.

For groupChromPeaks: a XCMSnExp object with the results of the peak grouping step (i.e. the features). These can be accessed with the featureDefinitions() method.

Slots

sampleGroups,ppm,absMz,minFraction,minSamples See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface which will eventually replace the group() methods. All of the settings to the algorithm can be passed with a MzClustParam object.

Calling groupChromPeaks on an XCMSnExp object will cause all eventually present previous correspondence results to be dropped.

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

The do_groupPeaks_mzClust() core API function and group.mzClust() for the old user interface.

featureDefinitions() and featureValues() for methods to access peak grouping results (i.e. the features).

XCMSnExp for the object containing the results of the peak grouping.

Other peak grouping methods: groupChromPeaks-density, groupChromPeaks-nearest, groupChromPeaks()

Examples

```
head(chromPeaks(fticr))
## Now create the MzClustParam parameter object: we're assuming here that
## both samples are from the same sample group.
p <- MzClustParam(sampleGroups = c(1, 1))
fticr <- groupChromPeaks(fticr, param = p)
## Get the definition of the features.
featureDefinitions(fticr)</pre>
```

groupChromPeaks-nearest

Peak grouping based on proximity in the mz-rt space

Description

This method is inspired by the grouping algorithm of mzMine (Katajamaa 2006) and performs correspondence based on proximity of peaks in the space spanned by retention time and mz values. The method creates first a *master peak list* consisting of all chromatographic peaks from the sample in which most peaks were identified, and starting from that, calculates distances to peaks from the sample with the next most number of peaks. If peaks are closer than the defined threshold they are grouped together.

The NearestPeaksParam class allows to specify all settings for the peak grouping based on the *nearest* algorithm. Instances should be created with the NearestPeaksParam constructor.

sampleGroups,sampleGroups<-: getter and setter for the sampleGroups slot of the object.

mzVsRtBalance,mzVsRtBalance<-: getter and setter for the mzVsRtBalance slot of the object.

absMz,absMz<-: getter and setter for the absMz slot of the object.

absRt,absRt<-: getter and setter for the absRt slot of the object.

kNN,kNN<-: getter and setter for the kNN slot of the object.

groupChromPeaks,XCMSnExp,NearestPeaksParam: performs peak grouping based on the proximity between chromatographic peaks from different samples in the mz-rt range.

The correspondence analysis can be performed on chromatographic peaks of any MS level (if present and if chromatographic peak detection has been performed for that MS level) defining features combining these peaks. The MS level can be selected with the parameter msLevel. By default, calling groupChromPeaks will remove any previous correspondence results. This can be disabled with add = TRUE, which will add newly defined features to already present feature definitions.

Usage

```
NearestPeaksParam(
   sampleGroups = numeric(),
   mzVsRtBalance = 10,
   absMz = 0.2,
   absRt = 15,
```

162

```
kNN = 10
)
## S4 method for signature 'NearestPeaksParam'
sampleGroups(object)
## S4 replacement method for signature 'NearestPeaksParam'
sampleGroups(object) <- value</pre>
## S4 method for signature 'NearestPeaksParam'
mzVsRtBalance(object)
## S4 replacement method for signature 'NearestPeaksParam'
mzVsRtBalance(object) <- value</pre>
## S4 method for signature 'NearestPeaksParam'
absMz(object)
## S4 replacement method for signature 'NearestPeaksParam'
absMz(object) <- value</pre>
## S4 method for signature 'NearestPeaksParam'
absRt(object)
## S4 replacement method for signature 'NearestPeaksParam'
absRt(object) <- value</pre>
## S4 method for signature 'NearestPeaksParam'
kNN(object)
```

S4 replacement method for signature 'NearestPeaksParam'
kNN(object) <- value</pre>

S4 method for signature 'XCMSnExp,NearestPeaksParam'
groupChromPeaks(object, param, msLevel = 1L, add = FALSE)

Arguments

sampleGroups	A vector of the same length than samples defining the sample group assignments (i.e. which samples belong to which sample group). This parameter is mandatory for the PeakDensityParam and has to be provided also if there is no sample grouping in the experiment (in which case all samples should be assigned to the same group).
mzVsRtBalance	numeric(1) representing the factor by which mz values are multiplied before calculating the (euclician) distance between two peaks.
absMz	numeric(1) maximum tolerated distance for mz values.
absRt	numeric(1) maximum tolerated distance for rt values.
kNN	numeric(1) representing the number of nearest neighbors to check.

object	For groupChromPeaks: an XCMSnExp object containing the results from a pre- vious chromatographic peak detection analysis (see findChromPeaks()).
	For all other methods: a `NearestPeaksParam` object.
value	The value for the slot.
param	A PeakDensityParam object containing all settings for the peak grouping algorithm.
msLevel	integer(1) defining the MS level. Currently only MS level 1 is supported.
add	logical(1) (default add = FALSE) allowing to perform an additional round of correspondence (e.g. on a different MS level) and add features to the already present feature definitions.

Value

The NearestPeaksParam function returns a NearestPeaksParam class instance with all of the settings specified for peak alignment based on peak proximity.

For groupChromPeaks: a XCMSnExp object with the results of the peak grouping/correspondence step (i.e. the mz-rt features). These can be accessed with the featureDefinitions() method.

Slots

sampleGroups, mzVsRtBalance, absMz, absRt, kNN See corresponding parameter above.

Note

These methods and classes are part of the updated and modernized xcms user interface. All of the settings to the algorithm can be passed with a NearestPeaksParam object.

References

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

See Also

The do_groupChromPeaks_nearest() core API function.

featureDefinitions() and featureValues() for methods to access peak grouping results (i.e. the features).

XCMSnExp for the object containing the results of the peak grouping.

Other peak grouping methods: groupChromPeaks-density, groupChromPeaks-mzClust, groupChromPeaks()

Examples

```
## Create a NearestPeaksParam object
p <- NearestPeaksParam(kNN = 3)
p</pre>
```

Load a test data set with detected peaks

groupFeatures-abundance-correlation

```
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahKO")</pre>
res <- faahko_sub
## Disable parallel processing for this example
register(SerialParam())
head(chromPeaks(res))
## The number of peaks identified per sample:
table(chromPeaks(res)[, "sample"])
## Performing the peak grouping
p <- NearestPeaksParam()</pre>
res <- groupChromPeaks(res, param = p)</pre>
## The results from the peak grouping:
featureDefinitions(res)
## Using the featureValues method to extract a matrix with the intensities of
## the features per sample.
head(featureValues(res, value = "into"))
## The process history:
processHistory(res)
```

groupFeatures-abundance-correlation

Compounding/feature grouping based on similarity of abundances across samples

Description

Features from the same originating compound are expected to have similar intensities across samples. This method this groups features based on similarity of abundances (i.e. *feature values*) across samples. See also AbundanceSimilarityParam() for additional information and details.

This help page lists parameters specific for xcms result objects (i.e. the XCMSnExp() object). Documentation of the parameters for the similarity calculation is available in the AbundanceSimilarityParam() help page in the MsFeatures package.

Usage

```
## S4 method for signature 'XCMSnExp,AbundanceSimilarityParam'
groupFeatures(
   object,
   param,
   msLevel = 1L,
   method = c("medret", "maxint", "sum"),
   value = "into",
```

```
intensity = "into",
filled = TRUE,
...
```

Arguments

object	XCMSnExp() object containing also correspondence results.
param	AbudanceSimilarityParam object with the settings for the method. See AbundanceSimilarityParam() for details on the grouping method and its parameters.
msLevel	integer(1) defining the MS level on which the features should be grouped.
method	<pre>character(1) passed to the featureValues call. See featureValues() for details. Defaults to method = "medret".</pre>
value	<pre>character(1) passed to the featureValues call. See featureValues() for details. Defaults to value = "into".</pre>
intensity	<pre>character(1) passed to the featureValues call. See featureValues() for details. Defaults to intensity = "into".</pre>
filled	logical(1) whether filled-in values should be included in the correlation anal- ysis. Defaults to filled = TRUE.
	additional parameters passed to the groupFeatures method for matrix.

Value

input XCMSnExp with feature group definitions added to a column "feature_group" in its featureDefinitions data frame.

Author(s)

Johannes Rainer

See Also

feature-grouping for a general overview.

Other feature grouping methods: groupFeatures-eic-similarity, groupFeatures-similar-rtime

Examples

```
library(MsFeatures)
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Group chromatographic peaks across samples</pre>
```

xodg <- groupChromPeaks(faahko_sub, param = PeakDensityParam(sampleGroups = rep(1, 3)))</pre>

166

Group features based on correlation of feature values (integrated ## peak area) across samples. Note that there are many missing values ## in the feature value which influence grouping of features in the present ## data set. xodg_grp <- groupFeatures(xodg, param = AbundanceSimilarityParam(threshold = 0.8)) table(featureDefinitions(xodg_grp)\$feature_group) ## Group based on the maximal peak intensity per feature xodg_grp <- groupFeatures(xodg, param = AbundanceSimilarityParam(threshold = 0.8, value = "maxo")) table(featureDefinitions(xodg_grp)\$feature_group)

groupFeatures-eic-similarity

Compounding/feature grouping based on similarity of extracted ion chromatograms

Description

Features from the same originating compound are expected to share their elution pattern (i.e. chromatographic peak shape) with it. Thus, this methods allows to group features based on similarity of their extracted ion chromatograms (EICs). The similarity calculation is performed separately for each sample with the similarity score being aggregated across samples for the final generation of the similarity matrix on which the grouping (considering parameter threshold) will be performed.

The compareChromatograms() function is used for similarity calculation which by default calculates the Pearson's correlation coefficient. The settings for compareChromatograms can be specified with parameters ALIGNFUN, ALIGNFUNARGS, FUN and FUNARGS. ALIGNFUN defaults to alignRt() and is the function used to *align* the chromatograms before comparison. ALIGNFUNARGS allows to specify additional arguments for the ALIGNFUN function. It defaults to ALIGNFUNARGS = list(tolerance = 0, method = "closest") which ensures that data points from the same spectrum (scan, i.e. with the same retention time) are compared between the EICs from the same sample. Parameter FUN defines the function to calculate the similarity score and defaults to FUN = cor and FUNARGS allows to pass additional arguments to this function (defaults to FUNARGS = list(use = "pairwise.complete.obs"). See also compareChromatograms() for more information.

The grouping of features based on the EIC similarity matrix is performed with the function specified with parameter groupFun which defaults to groupFun = groupSimilarityMatrix which groups all rows (features) in the similarity matrix with a similarity score larger than threshold into the same cluster. This creates clusters of features in which **all** features have a similarity score >= threshold with **any** other feature in that cluster. See groupSimilarityMatrix() for details. Additional parameters to that function can be passed with the ... argument.

This feature grouping should be called **after** an initial feature grouping by retention time (see SimilarRtimeParam()). The feature groups defined in columns "feature_group" of featureDefinitions(object) (for features matching msLevel) will be used and refined by this method. Features with a value of NA in featureDefinitions(object)\$feature_group will be skipped/not considered for feature grouping.

Usage

```
EicSimilarityParam(
  threshold = 0.9,
  n = 1,
  onlyPeak = TRUE,
  value = c("maxo", "into"),
  groupFun = groupSimilarityMatrix,
  ALIGNFUN = alignRt,
  ALIGNFUNARGS = list(tolerance = 0, method = "closest"),
  FUN = cor,
  FUNARGS = list(use = "pairwise.complete.obs"),
  ...
)
```

S4 method for signature 'XCMSnExp,EicSimilarityParam'
groupFeatures(object, param, msLevel = 1L)

Arguments

threshold	numeric(1) with the minimal required similarity score to group featues. This is passed to the groupFun function.
n	numeric(1) defining the total number of samples per feature group on which this similarity calculation should be performed. This value is rounded up to the next larger integer value.
onlyPeak	logical(1) whether the correlation should be performed only on the signals within the identified chromatographic peaks (onlyPeak = TRUE, default) or all the signal from the extracted ion chromatogram.
value	character(1) defining whether samples should be grouped based on the sum of the maximal peak intensity (value = "maxo", the default) or the integrated peak area (value = "into") for a feature.
groupFun	function defining the function to be used to group rows based on a pairwise similarity matrix. Defaults to groupSimilarityMatrix().
ALIGNFUN	<pre>function defining the function to be used to align chromatograms prior similar- ity calculation. Defaults to ALIGNFUN = alignRt. See alignRt() and compareChromatograms() for more information.</pre>
ALIGNFUNARGS	<pre>named list with arguments for ALIGNFUN. Defaults to ALIGNFUNARGS = list(tolerance = 0,method = "closest").</pre>
FUN	<pre>function defining the function to be used to calculate a similarity between (aligned) chromatograms. Defaults to FUN = cor. See cor() and compareChromatograms() for more information.</pre>
FUNARGS	named list with arguments for FUN. Defaults to FUN = list(use = "pairwise.complete.obs").
	for EicSimilarityParam: additional arguments to be passed to groupFun and featureChromatograms (such as expandRt to expand the retention time range of each feature).
object	XCMSnExp() object containing also correspondence results.

168

groupFeatures-eic-similarity

param	EicSimilarityParam object with the settings for the method.
msLevel	integer(1) defining the MS level on which the features should be grouped.

Value

input XCMSnExp with feature groups added (i.e. in column "feature_group" of its featureDefinitions data frame.

Note

While being possible to be performed on the full data set without prior feature grouping, this is not suggested for the following reasons: I) the selection of the top n samples with the highest signal for the *feature group* will be biased by very abundant compounds as this is performed on the full data set (i.e. the samples with the highest overall intensities are used for correlation of all features) and II) it is computationally much more expensive because a pairwise correlation between all features has to be performed.

It is also suggested to perform the correlation on a subset of samples per feature with the highest intensities of the peaks (for that feature) although it would also be possible to run the correlation on all samples by setting n equal to the total number of samples in the data set. EIC correlation should however be performed ideally on samples in which the original compound is highly abundant to avoid correlation of missing values or noisy peak shapes as much as possible.

By default also the signal which is outside identified chromatographic peaks is excluded from the correlation.

Author(s)

Johannes Rainer

See Also

feature-grouping for a general overview.

Other feature grouping methods: groupFeatures-abundance-correlation, groupFeatures-similar-rtime

Examples

```
library(MsFeatures)
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Group chromatographic peaks across samples
xodg <- groupChromPeaks(faahko_sub, param = PeakDensityParam(sampleGroups = rep(1, 3)))
## Performing a feature grouping based on EIC similarities on a single
## sample
xodg_grp <- groupFeatures(xodg, param = EicSimilarityParam(n = 1))</pre>
```

```
table(featureDefinitions(xodg_grp)$feature_group)
```

```
## Usually it is better to perform this correlation on pre-grouped features
## e.g. based on similar retention time.
xodg_grp <- groupFeatures(xodg, param = SimilarRtimeParam(diffRt = 4))
xodg_grp <- groupFeatures(xodg_grp, param = EicSimilarityParam(n = 1))</pre>
```

table(featureDefinitions(xodg_grp)\$feature_group)

groupFeatures-similar-rtime

Compounding/feature grouping based on similar retention times

Description

Group features based on similar retention time. This method is supposed to be used as an initial *crude* grouping of features based on the median retention time of all their chromatographic peaks. All features with a difference in their retention time which is <= parameter diffRt of the parameter object are grouped together. If a column "feature_group" is found in featureDefinitions() this is further sub-grouped by this method.

See MsFeatures::SimilarRtimeParam() in MsFeatures for more details.

Usage

```
## S4 method for signature 'XCMSnExp,SimilarRtimeParam'
groupFeatures(object, param, msLevel = 1L, ...)
```

Arguments

object	XCMSnExp() object containing also correspondence results.
param	SimilarRtimeParam object with the settings for the method. See MsFeatures::SimilarRtimeParam() for details and options.
msLevel	integer(1) defining the MS level on which the features should be grouped.
•••	passed to the groupFeatures function on numeric values.

Value

input XCMSnExp with feature groups added (i.e. in column "feature_group" of its featureDefinitions data frame.

Author(s)

Johannes Rainer

See Also

Other feature grouping methods: groupFeatures-abundance-correlation, groupFeatures-eic-similarity

Examples

```
library(MsFeatures)
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahKO")</pre>
## Disable parallel processing for this example
register(SerialParam())
## Group chromatographic peaks across samples
xodg <- groupChromPeaks(faahko_sub, param = PeakDensityParam(sampleGroups = rep(1, 3)))</pre>
## Group features based on similar retention time (i.e. difference <= 2 seconds)</pre>
xodg_grp <- groupFeatures(xodg, param = SimilarRtimeParam(diffRt = 2))</pre>
## Feature grouping get added to the featureDefinitions in column "feature_group"
head(featureDefinitions(xodg_grp)$feature_group)
table(featureDefinitions(xodg_grp)$feature_group)
length(unique(featureDefinitions(xodg_grp)$feature_group))
## Using an alternative groupiing method that creates larger groups
xodg_grp <- groupFeatures(xodg,</pre>
    param = SimilarRtimeParam(diffRt = 2, groupFun = MsCoreUtils::group))
```

length(unique(featureDefinitions(xodg_grp)\$feature_group))

groupnames, XCMSnExp-method

Generate unique group (feature) names based on mass and retention time

Description

groupnames generates names for the identified features from the correspondence analysis based in their mass and retention time. This generates feature names that are equivalent to the group names of the *old* user interface (aka xcms1).

Usage

```
## S4 method for signature 'XCMSnExp'
groupnames(object, mzdec = 0, rtdec = 0, template = NULL)
```

Arguments

object	XCMSnExp object containing correspondence results.
mzdec	integer(1) with the number of decimal places to use for m/z (defaults to 0).

	integer(1) with the number of decimal places to use for the retention time (defaults to \emptyset).
template	character with existing group names whose format should be emulated.

Value

character with unique names for each feature in object. The format is M(m/z)T(time in seconds).

See Also

XCMSnExp.

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)

See Also

xcmsSet-class, xcmsEIC-class

groupOverlaps

Description

groupOverlaps identifies overlapping ranges in the input data and groups them by returning their indices in xmin xmax.

Usage

groupOverlaps(xmin, xmax)

Arguments

xmin	numeric (same length than xmax) with the lower boundary of the range.
xmax	numeric (same length than xmin) with the upper boundary of the range.

Value

list with the indices of grouped elements.

Author(s)

Johannes Rainer

Examples

x <- c(2, 12, 34.2, 12.4) y <- c(3, 16, 35, 36)

groupOverlaps(x, y)

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

See Also

xcmsSet-class

highlightChromPeaks	Add definition of chromatographic peaks to an extracted ch	ro-
	matogram plot	

Description

The highlightChromPeaks function adds chromatographic peak definitions to an existing plot, such as one created by the plot method on a Chromatogram or MChromatograms object.

Usage

```
highlightChromPeaks(
    x,
    rt,
    mz,
    peakIds = character(),
    border = rep("00000040", length(fileNames(x))),
    lwd = 1,
    col = NA,
    type = c("rect", "point", "polygon"),
    whichPeaks = c("any", "within", "apex_within"),
    ...
)
```

Arguments

x	For highlightChromPeaks: XCMSnExp object with the detected peaks.
rt	For highlightChromPeaks: numeric(2) with the retention time range from which peaks should be extracted and plotted.
mz	numeric(2) with the mz range from which the peaks should be extracted and plotted.
peakIds	character defining the IDs (i.e. rownames of the peak in the chromPeaks table) of the chromatographic peaks to be highlighted in a plot.
border	colors to be used to color the border of the rectangles/peaks. Has to be equal to the number of samples in x.
lwd	numeric(1) defining the width of the line/border.
col	For highlightChromPeaks: color to be used to fill the rectangle (if type = "rect") or the peak (for type = "polygon").
type	the plotting type. See plot in base grapics for more details. For highlightChromPeaks: character(1) defining how the peak should be highlighted: type = "rect" draws a rectangle representing the peak definition, type = "point" indicates a chromatographic peak with a single point at the position of the peak's "rt" and "maxo" and type = "polygon" will highlight the peak shape. For type = "polygon" the color of the border and area can be defined with parameters "border" and "col", respectively.
whichPeaks	character(1) specifying how peaks are called to be located within the region defined by mz and rt. Can be one of "any", "within", and "apex_within" for all peaks that are even partially overlapping the region, peaks that are completely within the region, and peaks for which the apex is within the region. This pa- rameter is passed to the type argument of the chromPeaks function. See related documentation for more information and examples.
	additional parameters to the matplot or plot function.

Author(s)

Johannes Rainer

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Disable parallel processing for this example
register(SerialParam())
## Extract the ion chromatogram for one chromatographic peak in the data.
chrs <- chromatogram(faahko_sub, rt = c(2700, 2900), mz = 335)
plot(chrs)</pre>
```

image-methods Plot log intensity image of a xcmsRaw ob	vject
--	-------

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

imputeLinInterpol Impute values for empty elements in a vector using linear interpolation

Description

This function provides missing value imputation based on linear interpolation and resembles some of the functionality of the profBinLin and profBinLinBase functions deprecated from version 1.51 on.

Usage

```
imputeLinInterpol(
    x,
    baseValue,
    method = "lin",
    distance = 1L,
    noInterpolAtEnds = FALSE
)
```

Arguments

х	A numeric vector with eventual missing (NA) values.
baseValue	The base value to which empty elements should be set. This is only considered for method = "linbase" and corresponds to the profBinLinBase's baselevel argument.
method	One of "none", "lin" or "linbase".
distance	For method = "linbase": number of non-empty neighboring element of an empty element that should be considered for linear interpolation. See details section for more information.
noInterpolAtEnds	
	For method = "lin": Logical indicating whether linear interpolation should also be performed at the ends of the data vector (i.e. if missing values are present at the beginning or the end of the vector).

Details

Values for NAs in input vector x can be imputed using methods "lin" and "linbase":

impute = "lin" uses simple linear imputation to derive a value for an empty element in input vector x from its neighboring non-empty elements. This method is equivalent to the linear interpolation in the profBinLin method. Whether interpolation is performed if missing values are present at the beginning and end of x can be set with argument noInterpolAtEnds. By default interpolation is also performed at the ends interpolating from 0 at the beginning and towards 0 at the end. For noInterpolAtEnds = TRUE no interpolation is performed at both ends replacing the missing values at the beginning and/or the end of x with 0. impute = "linbase" uses linear interpolation to impute values for empty elements within a userdefinable proximity to non-empty elements and setting the element's value to the baseValue otherwise. The default for the baseValue is half of the smallest value in x (NAs being removed). Whether linear interpolation based imputation is performed for a missing value depends on the distance argument. Interpolation is only performed if one of the next distance closest neighbors to the current empty element has a value other than NA. No interpolation takes place for distance = 0, while distance = 1 means that the value for an empty element is interpolated from directly adjacent non-empty elements while, if the next neighbors of the current empty element are also NA, it's vale is set to baseValue. This corresponds to the linear interpolation performed by the profBinLinBase method. For more details see examples below.

Value

A numeric vector with empty values imputed based on the selected method.

Author(s)

Johannes Rainer

Examples

```
#######
## Impute missing values by linearly interpolating from neighboring
## non-empty elements
x <- c(3, NA, 1, 2, NA, NA, 4, NA, NA, NA, 3, NA, NA, NA, NA, 2)
imputeLinInterpol(x, method = "lin")
## visualize the interpolation:
plot(x = 1:length(x), y = x)
points(x = 1:length(x), y = imputeLinInterpol(x, method = "lin"), type = "l", col = "grey")
## If the first or last elements are NA, interpolation is performed from 0
## to the first non-empty element.
x <- c(NA, 2, 1, 4, NA)
imputeLinInterpol(x, method = "lin")
## visualize the interpolation:
plot(x = 1:length(x), y = x)
points(x = 1:length(x), y = x)
points(x = 1:length(x), y = imputeLinInterpol(x, method = "lin"), type = "l", col = "grey")</pre>
```

```
## If noInterpolAtEnds is TRUE no interpolation is performed at both ends
imputeLinInterpol(x, method = "lin", noInterpolAtEnds = TRUE)
```

######

method = "linbase"
"linbase" performs imputation by interpolation for empty elements based on
'distance' adjacent non-empty elements, setting all remaining empty elements
to the baseValue
x <- c(3, NA, 1, 2, NA, NA, 4, NA, NA, NA, 3, NA, NA, NA, NA, 2)
Setting distance = 0 skips imputation by linear interpolation
imputeLinInterpol(x, method = "linbase", distance = 0)</pre>

With distance = 1 for all empty elements next to a non-empty element the value
is imputed by linear interpolation.

imputeRowMin

```
xInt <- imputeLinInterpol(x, method = "linbase", distance = 1L)
xInt
plot(x = 1:length(x), y = x, ylim = c(0, max(x, na.rm = TRUE)))
points(x = 1:length(x), y = xInt, type = "1", col = "grey")
## Setting distance = 2L would cause that for all empty elements for which the
## distance to the next non-empty element is <= 2 the value is imputed by
## linear interpolation:
xInt <- imputeLinInterpol(x, method = "linbase", distance = 2L)
xInt
plot(x = 1:length(x), y = x, ylim = c(0, max(x, na.rm = TRUE)))
points(x = 1:length(x), y = xInt, type = "1", col = "grey")</pre>
```

imputeRowMin

Replace missing values with a proportion of the row minimum

Description

imputeRowMin imputes missing values in x by replacing NAs in each row with a proportion of the minimal value for that row (i.e. min_fraction * min(x[i,])).

Usage

```
imputeRowMin(x, min_fraction = 1/2)
```

Arguments

х	matrix with abundances, rows being features/metabolites and columns samples.
min_fraction	numeric(1) with the fraction of the row minimum that should be used to replace NA values in that row.

Author(s)

Johannes Rainer

See Also

imputeLCMD package for more left censored imputation functions.

Other imputation functions: imputeRowMinRand()

Examples

```
library(faahKO)
data("faahko")
xset <- group(faahko)
mat <- groupval(xset, value = "into")
mat_imp <- imputeRowMin(mat)
head(mat)
head(mat_imp)
## Replace with 1/8 of the row mimimum
head(imputeRowMin(mat, min_fraction = 1/8))</pre>
```

imputeRowMinRand

Impute missing values with random numbers based on the row minimum

Description

Replace missing values with random numbers. When using the method = "mean_sd", random numbers will be generated from a normal distribution based on (a fraction of) the row min and a standard deviation estimated from the linear relationship between row standard deviation and mean of the full data set. Parameter sd_fraction allows to further reduce the estimated standard deviation. When using the method method = "from_to", random numbers between 2 specific values will be generated.

Usage

```
imputeRowMinRand(
    x,
    method = c("mean_sd", "from_to"),
    min_fraction = 1/2,
    min_fraction_from = 1/1000,
    sd_fraction = 1,
    abs = TRUE
)
```

Arguments

х	matrix with abundances, rows being features/metabolites and columns samples.
method	<pre>method character(1) defining the imputation method. See description for de- tails. Defaults to method = "mean_sd".</pre>
min_fraction	numeric(1) with the fraction of the row minimum that should be used to replace NA values in that row in case that mean_sd method is specified. When using from_to method, this value will be the one used to calculate the maximum value for replace NA values in that row.

180

<pre>min_fraction_from</pre>

	numeric(1) with the fraction of the row minimum that should be used to calcu- late the minimum value for replace NA values in that row. This parameter is used only in case that from_to method is specified.
sd_fraction	numeric(1) factor to reduce the estimated standard deviation. This parameter is used only in case that mean_sd method is specified.
abs	logical(1) to force imputed values to be strictly positive.

Details

For method **mean_sd**, imputed values are taken from a normal distribution with mean being a user defined fraction of the row minimum and the standard deviation estimated for that mean based on the linear relationship between row standard deviations and row means in the full matrix x.

To largely avoid imputed values being negative or larger than the *real* values, the standard deviation for the random number generation is estimated ignoring the intercept of the linear model estimating the relationship between standard deviation and mean. If abs = TRUE NA values are replaced with the absolute value of the random values.

For method **from_to**, imputed values are taken between 2 user defined fractions of the row minimum.

Author(s)

Johannes Rainer, Mar Garcia-Aloy

See Also

imputeLCMD package for more left censored imputation functions.
Other imputation functions: imputeRowMin()

Examples

```
library(faahKO)
data("faahko")
xset <- group(faahko)
mat <- groupval(xset, value = "into")
## Estimate the relationship between row sd and mean. The standard deviation
## of the random distribution is estimated on this relationship.
mns <- rowMeans(mat, na.rm = TRUE)
sds <- apply(mat, mARGIN = 1, sd, na.rm = TRUE)
plot(mns, sds)
abline(lm(sds ~ mns))
mat_imp_meansd <- imputeRowMinRand(mat, method = "mean_sd")
mat_imp_fromto <- imputeRowMinRand(mat, method = "from_to")
head(mat)
head(mat_imp_meansd)
head(mat_imp_fromto)
```

Description

isolationWindowTargetMz extracts the isolation window target m/z definition for each spectrum in object.

Usage

S4 method for signature 'OnDiskMSnExp'
isolationWindowTargetMz(object)

Arguments

object OnDiskMSnExp object.

Value

a numeric of length equal to the number of spectra in object with the isolation window target m/z or NA if not specified/available.

Author(s)

Johannes Rainer

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.

loadRaw-methods

Methods

x = "xcmsRaw" levelplot(x,log=TRUE,col.regions=colorRampPalette(brewer.pal(9,"YlOrRd"))(256),...)

x = "xcmsSet" levelplot(x,log=TRUE,col.regions=colorRampPalette(brewer.pal(9,"YlOrRd"))(256),rt="ra

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 the following components:

Numeric vector with acquisition time (in seconds) for each scan
Numeric vector with Total Ion Count for each scan
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]
Concatenated vector of m/z values for all scans
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

manualChromPeaks Manual peak integration and feature definition

Description

The manualChromPeaks function allows to manually define chromatographic peaks which are added to the object's chromPeaks matrix. In contrast to findChromPeaks(), no *peak detection* is performed (e.g. using an algorithm such as *centWave*) but the peak is added as defined by the user. Note that a peak will not be added if no signal (intensity) was found in a sample within the provided boundaries.

Because chromatographic peaks are added to eventually previously identified peaks, it is suggested to run refineChromPeaks() with the MergeNeighboringPeaksParam() approach to merge potentially overlapping peaks.

The manualFeatures function allows to manually group identified chromatographic peaks into features by providing their index in the object's chromPeaks matrix.

Usage

```
manualChromPeaks(
   object,
   chromPeaks = matrix(),
   samples = seq_along(fileNames(object)),
   BPPARAM = bpparam(),
   msLevel = 1L
)
```

manualFeatures(object, peakIdx = list(), msLevel = 1L)

Arguments

object XCMSnExp or OnDiskMSnExp object. chromPeaks matrix defining the boundaries of the chromatographic peaks, one row per chromatographic peak, columns "mzmin", "mzmax", "rtmin" and "rtmax" defining the m/z and retention time region of each peak.

medianFilter

samples	optional integer to select samples in which the peak integration should be per- formed. By default performed in all samples.
BPPARAM	parallel processing settings (see <pre>bpparam()</pre> for details).
msLevel	integer(1) defining the MS level in which peak integration should be performed.
peakIdx	for nabbyakFeatyres: list of integer vectors with the indices of chromato- graphic peaks in the object's chromPeaks matrix that should be grouped into features.

Value

XCMSnExp with the manually added chromatographic peaks or features.

Author(s)

Johannes Rainer

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>
```

MergeNeighboringPeaksParam

Merge neighboring and overlapping chromatographic peaks

Description

Peak detection sometimes fails to identify a chromatographic peak correctly, especially for broad peaks and if the peak shape is irregular (mostly for HILIC data). In such cases several smaller peaks are reported. Also, peak detection can result in partially or completely overlapping peaks. To reduce such peak detection artifacts, this function merges chromatographic peaks which are overlapping or close in rt and m/z dimension considering also the measured signal intensities in the region between them.

Chromatographic peaks are first expanded in m/z and retention time dimension (based on parameters expandMz, ppm and expandRt) and subsequently grouped into sets of merge candidates if they are (after expansion) overlapping in both m/z and rt (within the same sample). Candidate peaks are merged if the average intensity of the 3 data points in the middle position between them (i.e. at half the distance between "rtmax" of the first and "rtmin" of the second peak) is larger than a certain proportion (minProp) of the smaller maximal intensity ("maxo") of both peaks. In cases in which this calculated mid point is **not** located between the apexes of the two peaks (e.g. if the peaks are largely overlapping) the average signal intensity at half way between the apexes is used instead. Candidate peaks are not joined if all 3 data points between them have NA intensities. The joined peaks get the "mz", "rt", "sn" and "maxo" values from the peak with the largest signal ("maxo") as well as its row in the metadata data frame of the peak (chromPeakData). The "rtmin", "rtmax" of the merged peaks are updated and "into" is recalculated based on all the signal between "rtmin" and "rtmax" of the new merged peak. See details for information on the "mzmin" and "mzmax" values of the merged peaks.

Usage

```
MergeNeighboringPeaksParam(
    expandRt = 2,
    expandMz = 0,
    ppm = 10,
    minProp = 0.75
)
## S4 method for signature 'XCMSnExp,MergeNeighboringPeaksParam'
refineChromPeaks(
    object,
    param = MergeNeighboringPeaksParam(),
    msLevel = 1L,
    BPPARAM = bpparam()
)
```

expandRt	numeric(1) defining by how many seconds the retention time window is expanded on both sides to check for overlapping peaks.
expandMz	numeric(1) constant value by which the m/z range of each chromatographic peak is expanded (on both sides!) to check for overlapping peaks.
ppm	numeric(1) defining a m/z relative value (in parts per million) by which the m/z range of each chromatographic peak is expanded to check for overlapping peaks.
minProp	numeric(1) between 0 and 1 representing the proportion of intensity to be re- quired for peaks to be joined. See description for more details. The default (minProp = 0.75) means that peaks are only joined if the signal half way be- tween then is larger 75% of the smallest of the two peak's "maxo" (maximal intensity at peak apex).
object	XCMSnExp object with identified chromatographic peaks.
param	MergeNeighboringPeaksParam object defining the settings for the method.
msLevel	integer defining for which MS level(s) the chromatographic peaks should be merged.
BPPARAM	parameter object to set up parallel processing. Uses the default parallel process- ing setup returned by bpparam(). See bpparam() for details and examples.

Details

For each set of candidate peaks an ion chromatogram is extracted using the range of retention times and m/z values of these peaks. The m/z range for the extracted ion chromatogram is expanded by expandMz and ppm (on both sides) to reduce the possibility of missing signal intensities between candidate peaks (variance of measured m/z values for lower intensities is larger than for higher intensities and thus data points not being part of identified chromatographic peaks tend to have m/z values outside of the m/z range of the candidate peaks - especially for ToF instruments). This also ensures that all data points from the same ion are considered for the peak integration of merged peaks. The smallest and largest m/z value of all data points used in the peak integration of the merged peak are used as the merged peak's m/z range (i.e. columns "mzmin" and "mzmax").

Value

XCMSnExp object with chromatographic peaks matching the defined conditions being merged.

Note

Note that **each** peak gets expanded by expandMz and expandRt, thus peaks differing by 2 * expandMz (or expandRt) will be identified as *overlapping*. As an example: m/z max of one peak is 12.2, m/z min of another one is 12.4, if expandMz = 0.1 the m/z max of the first peak will be 12.3 and the m/z min of the second one 12.3, thus both are considered overlapping.

refineChromPeaks methods will always remove feature definitions, because a call to this method can change or remove identified chromatographic peaks, which may be part of features.

Merging of chromatographic peaks is performed along the retention time axis, i.e. candidate peaks are first ordered by their "rtmin" value. The signals at half way between the first and the second

candidate peak are then compared to the smallest "maxo" of both and the two peaks are then merged if the average signal between the peaks is larger minProp. For merging any additional peak in a candidate peak list the "maxo" of that peak and the newly merged peak are considered.

Author(s)

Johannes Rainer, Mar Garcia-Aloy

See Also

Other chromatographic peak refinement methods: CleanPeaksParam, FilterIntensityParam

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahKO")</pre>
## Disable parallel processing for this example
register(SerialParam())
## Subset to a single file
xd <- filterFile(faahko_sub, file = 1)</pre>
## Example of a split peak that will be merged
mzr <- 305.1 + c(-0.01, 0.01)
chr <- chromatogram(xd, mz = mzr, rt = c(2700, 3700))
plot(chr)
## Combine the peaks
res <- refineChromPeaks(xd, param = MergeNeighboringPeaksParam(expandRt = 4))</pre>
chr_res <- chromatogram(res, mz = mzr, rt = c(2700, 3700))</pre>
plot(chr_res)
## Example of a peak that was not merged, because the signal between them
## is lower than the cut-off minProp
mzr <- 496.2 + c(-0.01, 0.01)
chr <- chromatogram(xd, mz = mzr, rt = c(3200, 3500))</pre>
plot(chr)
chr_res <- chromatogram(res, mz = mzr, rt = c(3200, 3500))
plot(chr_res)
```

```
msn2xcmsRaw
```

Copy MSn data in an xcmsRaw to the MS slots

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.

overlappingFeatures

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>
```

overlappingFeatures Identify overlapping features

Description

overlappingFeatures identifies features that are overlapping or close in the m/z - rt space.

Usage

```
overlappingFeatures(x, expandMz = 0, expandRt = 0, ppm = 0)
```

х	XCMSnExp with the features.
expandMz	numeric(1) with the value to expand each feature (on each side) in m/z di- mension before identifying overlapping features. The resulting "mzmin" for the feature is thus mzmin -expandMz and the "mzmax" mzmax + expandMz.
expandRt	numeric(1) with the value to expand each feature (on each side) in retention time dimension before identifying overlapping features. The resulting "rtmin" for the feature is thus rtmin -expandRt and the "rtmax" rtmax + expandRt.
ppm	numeric(1) to grow the m/z width of the feature by a relative value: mzmin -mzmin * ppm / 2e6, mzmax + mzmax * ppm / 2e6. Each feature is thus expanded in m/z dimension by ppm/2 on each side before identifying overlapping features.

Value

list with indices of features (in featureDefinitions()) that are overlapping.

Author(s)

Johannes Rainer

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")
## Disable parallel processing for this example
register(SerialParam())
## Correspondence analysis
xdata <- groupChromPeaks(faahko_sub, param = PeakDensityParam(sampleGroups = c(1, 1, 1)))
## Identify overlapping features
overlappingFeatures(xdata)
## Identify features that are separated on retention time by less than
## 2 minutes
overlappingFeatures(xdata, expandRt = 60)
```

peakPlots-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.

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

peaksWithCentWave Identify peaks in chromatographic data using centWave

Description

peaksWithCentWave identifies (chromatographic) peaks in purely chromatographic data, i.e. based on intensity and retention time values without m/z values.

Usage

```
peaksWithCentWave(
    int,
    rt,
    peakwidth = c(20, 50),
    snthresh = 10,
    prefilter = c(3, 100),
    integrate = 1,
    fitgauss = FALSE,
    noise = 0,
    verboseColumns = FALSE,
    firstBaselineCheck = TRUE,
    extendLengthMSW = FALSE,
    ...
)
```

int	numeric with intensity values.	
rt	numeric with the retention time for the intensities. Length has to be equal to length(int).	
peakwidth	numeric(2) with the lower and upper bound of the expected peak width.	
snthresh	numeric(1) defining the signal to noise ratio cutoff. Peaks with a signal to noise ratio < snthresh are omitted.	
prefilter	numeric(2) (c(k,I)): only regions of interest with at least k centroids with signal \geq I are returned in the first step.	
integrate	numeric(1), integration method. For integrate = 1 peak limits are found through descending on the mexican hat filtered data, for integrate = 2 the de- scend is done on the real data. The latter method is more accurate but prone to noise, while the former is more robust, but less exact.	
fitgauss	logical(1) whether or not a Gaussian should be fitted to each peak.	
noise	numeric(1) defining the minimum required intensity for centroids to be con- sidered in the first analysis step (definition of the <i>regions of interest</i>).	
verboseColumns	logical(1): whether additional peak meta data columns should be returned.	
firstBaselineCheck		
	logical(1). If TRUE continuous data within regions of interest is checked to be above the first baseline.	
extendLengthMSW		
	logical(1). If TRUE the "open" method of EIC extension is used, rather than the default "reflect" method.	
	currently ignored.	

Details

The method uses the same algorithm for the peak detection than centWave, employs however a different approach to identify the initial regions in which the peak detection is performed (i.e. the *regions of interest* ROI). The method first identifies all local maxima in the chromatographic data and defines the corresponding positions +/- peakwidth[2] as the ROIs. Noise estimation bases also on these ROIs and can thus be different from centWave resulting in different signal to noise ratios.

Value

A matrix, each row representing an identified chromatographic peak, with columns:

- "rt": retention time of the peak's midpoint (time of the maximum signal).
- "rtmin": minimum retention time of the peak.
- "rtmax": maximum retention time of the peak.
- "into": integrated (original) intensity of the peak.
- "intb": per-peak baseline corrected integrated peak intensity.
- "maxo": maximum (original) intensity of the peak.

peaksWithCentWave

• "sn": signal to noise ratio of the peak defined as (maxo -baseline)/sd with sd being the standard deviation of the local chromatographic noise.

Additional columns for verboseColumns = TRUE:

- "mu": gaussian parameter mu.
- "sigma": gaussian parameter sigma.
- "h": gaussian parameter h.
- "f": region number of the m/z ROI where the peak was localized.
- "dppm": m/z deviation of mass trace across scans in ppm (always NA).
- "scale": scale on which the peak was localized.
- "scpos": peak position found by wavelet analysis (index in int).
- "scmin": left peak limit found by wavelet analysis (index in int).
- "scmax": right peak limit found by wavelet analysis (index in int).

Author(s)

Johannes Rainer

See Also

centWave for a detailed description of the peak detection method.

Other peak detection functions for chromatographic data: peaksWithMatchedFilter()

Examples

```
## Reading a file
library(xcms)
od <- readMSData(system.file("cdf/KO/ko15.CDF", package = "faahKO"),</pre>
    mode = "onDisk")
## Extract chromatographic data for a small m/z range
mzr <- c(272.1, 272.2)</pre>
chr <- chromatogram(od, mz = mzr, rt = c(3000, 3300))[1, 1]
int <- intensity(chr)</pre>
rt <- rtime(chr)</pre>
## Plot the region
plot(chr, type = "h")
## Identify peaks in the chromatographic data
pks <- peaksWithCentWave(intensity(chr), rtime(chr))</pre>
pks
## Highlight the peaks
rect(xleft = pks[, "rtmin"], xright = pks[, "rtmax"],
    ybottom = rep(0, nrow(pks)), ytop = pks[, "maxo"], col = "#ff000040",
    border = "#00000040")
```

peaksWithMatchedFilter

Identify peaks in chromatographic data using matchedFilter

Description

The function performs peak detection using the matchedFilter algorithm on chromatographic data (i.e. with only intensities and retention time).

Usage

```
peaksWithMatchedFilter(
    int,
    rt,
    fwhm = 30,
    sigma = fwhm/2.3548,
    max = 20,
    snthresh = 10,
    ...
)
```

Arguments

int	numeric with intensity values.
rt	numeric with the retention time for the intensities. Length has to be equal to length(int).
fwhm	numeric(1) specifying the full width at half maximum of matched filtration gaussian model peak. Only used to calculate the actual sigma, see below.
sigma	numeric(1) specifying the standard deviation (width) of the matched filtration model peak.
max	numeric(1) with the maximal number of peaks that are expected/ will bbe de- tected in the data
snthresh	numeric(1) defining the signal to noise cut-off to be used in the peak detection step.
	currently ignored.

Value

A matrix, each row representing an identified chromatographic peak, with columns:

- "rt": retention time of the peak's midpoint (time of the maximum signal).
- "rtmin": minimum retention time of the peak.
- "rtmax": maximum retention time of the peak.
- "into": integrated (original) intensity of the peak.

peakTable-methods

- "intf": integrated intensity of the filtered peak.
- "maxo": maximum (original) intensity of the peak.
- "maxf"" maximum intensity of the filtered peak.
- "sn": signal to noise ratio of the peak.

Author(s)

Johannes Rainer

See Also

matchedFilter for a detailed description of the peak detection method.

Other peak detection functions for chromatographic data: peaksWithCentWave()

Examples

```
## Load the test file
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")
## Subset to one file and drop identified chromatographic peaks
data <- dropChromPeaks(filterFile(faahko_sub, 1))
## Extract chromatographic data for a small m/z range
chr <- chromatogram(data, mz = c(272.1, 272.3), rt = c(3000, 3200))[1, 1]
pks <- peaksWithMatchedFilter(intensity(chr), rtime(chr))
pks
## Plotting the data
plot(rtime(chr), intensity(chr), type = "h")
rect(xleft = pks[, "rtmin"], xright = pks[, "rtmax"], ybottom = c(0, 0),
    ytop = pks[, "maxo"], border = "red")
```

peakTable-methods Create report of aligned peak intensities

Description

Create a report showing all aligned peaks.

Arguments

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)

phenoDataFromPaths Derive experimental design from file paths

Description

The phenoDataFromPaths function builds a data.frame representing the experimental design from the folder structure in which the files of the experiment are located.

Usage

```
phenoDataFromPaths(paths)
```

Arguments

paths

character representing the file names (including the full path) of the experiment's files.

Note

This function is used by the *old* xcmsSet function to guess the experimental design (i.e. group assignment of the files) from the folders in which the files of the experiment can be found.

Examples

```
## List the files available in the faahKO package
base_dir <- system.file("cdf", package = "faahKO")
cdf_files <- list.files(base_dir, recursive = TRUE, full.names = TRUE)</pre>
```

plot.xcmsEIC

Plot extracted ion chromatograms from multiple files

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

Author(s)

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

See Also

xcmsEIC-class, png, pdf, postscript,

plotAdjustedRtime Visualization of alignment results

Description

Plot the difference between the adjusted and the raw retention time (y-axis) for each file along the (adjusted or raw) retention time (x-axis). If alignment was performed using the adjustRtime-peakGroups method, also the features (peak groups) used for the alignment are shown.

plotAdjustedRtime

Usage

```
plotAdjustedRtime(
    object,
    col = "#0000080",
    lty = 1,
    lwd = 1,
    type = "l",
    adjustedRtime = TRUE,
    xlab = ifelse(adjustedRtime, yes = expression(rt[adj]), no = expression(rt[raw])),
    ylab = expression(rt[adj] - rt[raw]),
    peakGroupsCol = "#00000060",
    peakGroupsCol = "#0000060",
    peakGroupsLty = 3,
    ylim,
    ....
)
```

Arguments

object	A XCMSnExp object with the alignment results.
col	colors to be used for the lines corresponding to the individual samples.
lty	line type to be used for the lines of the individual samples.
lwd	line width to be used for the lines of the individual samples.
type	plot type to be used. See help on the par function for supported values.
adjustedRtime	logical(1) whether adjusted or raw retention times should be shown on the x-axis.
xlab	the label for the x-axis.
ylab	the label for the y-axis.
peakGroupsCol	color to be used for the peak groups (only used if alignment was performed using the adjustRtime-peakGroups method.
peakGroupsPch	point character (pch) to be used for the peak groups (only used if alignment was performed using the adjustRtime-peakGroups method.
peakGroupsLty	line type (lty) to be used to connect points for each peak groups (only used if alignment was performed using the adjustRtime-peakGroups method.
ylim	optional numeric(2) with the upper and lower limits on the y-axis.
	Additional arguments to be passed down to the plot function.

Author(s)

Johannes Rainer

See Also

adjustRtime for all retention time correction/ alignment methods.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")</pre>
## Disable parallel processing for this example
register(SerialParam())
## Performing the peak grouping using the "peak density" method.
p <- PeakDensityParam(sampleGroups = c(1, 1, 1))</pre>
res <- groupChromPeaks(faahko_sub, param = p)</pre>
## Perform the retention time adjustment using peak groups found in both
## files.
fgp <- PeakGroupsParam(minFraction = 1)</pre>
res <- adjustRtime(res, param = fgp)</pre>
## Visualize the impact of the alignment.
plotAdjustedRtime(res, adjusted = FALSE)
grid()
```

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

See Also

xcmsRaw-class

plotChromatogramsOverlay

Plot multiple chromatograms into the same plot

Description

plotOverlay draws chromatographic peak data from multiple (different) extracted ion chromatograms (EICs) into the same plot. This allows to directly compare the peak shape of these EICs in the same sample. In contrast to the plot function for MChromatograms() object, which draws the data from the same EIC across multiple samples in the same plot, this function draws the different EICs from the same sample into the same plot.

If plotChromatogramsOverlay is called on a XChromatograms object any present chromatographic peaks will also be highlighted/drawn depending on the parameters peakType, peakCol, peakBg and peakPch (see also help on the plot function for XChromatogram() object for details).

Usage

```
## S4 method for signature 'MChromatograms'
plotChromatogramsOverlay(
  object,
  col = "#00000060",
  type = "1",
 main = NULL,
 xlab = "rtime",
 ylab = "intensity",
 xlim = numeric(),
 ylim = numeric(),
  stacked = 0,
  transform = identity,
)
## S4 method for signature 'XChromatograms'
plotChromatogramsOverlay(
  object,
  col = "#00000060",
  type = "1",
 main = NULL,
```

```
xlab = "rtime",
ylab = "intensity",
xlim = numeric(),
peakType = c("polygon", "point", "rectangle", "none"),
peakBg = NULL,
peakCol = NULL,
peakPch = 1,
stacked = 0,
transform = identity,
...
```

object	MChromatograms() or XChromatograms() object.
col	definition of the color in which the chromatograms should be drawn. Can be of length 1 or equal to nrow(object) to plot each overlayed chromatogram in a different color.
type	character(1) defing the type of the plot. By default (type = "1") each chro- matogram is drawn as a line.
main	optional title of the plot. If not defined, the range of m/z values is used.
xlab	character(1) defining the x-axis label.
ylab	character(1) defining the y-axis label.
xlim	optional numeric(2) defining the x-axis limits.
ylim	optional numeric(2) defining the y-axis limits.
stacked	numeric(1) defining the part (proportion) of the y-axis to use to <i>stack</i> EICs depending on their m/z values. If stacked = 0 (the default) no stacking is performed. With stacked = 1 half of the y-axis is used for stacking and half for the intensity y-axis (i.e. the ratio between stacking and intensity y-axis is 1:1). Note that if stacking is different from 0 no y-axis and label are drawn.
transform	<pre>function to transform the intensity values before plotting. Defaults to transform = identity which plots the data as it is. With transform = log10 intensity val- ues would be log10 transformed before plotting.</pre>
	optional arguments to be passed to the plotting functions (see help on the base R plot function.
peakType	if object is a XChromatograms object: how chromatographic peaks should be drawn: peakType = "polygon" (the default): label the full chromatographic peak area, peakType = "rectangle": indicate the chromatographic peak by a rectangle and peakType = "point": label the chromatographic peaks' apex po- sition with a point.
peakBg	if object is a XChromatograms object: definition of background color(s) for each chromatographic peak. Has to be either of length 1 or equal to the number of peaks in object. If not specified, the peak will be drawn in the color defined by col.

peakCol	if object is a XChromatograms object: definition of color(s) for each chromato-
	graphic peak. Has to be either of length 1 or equal to the number of peaks in
	object. If not specified, the peak will be drawn in the color defined by col.
peakPch	if object is a XChromatograms object: <i>point character</i> to be used to label the apex position of the chromatographic peak if peakType = "point".

Value

silently returns a list (length equal to ncol(object) of numeric (length equal to nrow(object)) with the y position of each EIC.

Author(s)

Johannes Rainer

Examples

```
## Load preprocessed data and extract EICs for some features.
library(xcms)
data(xdata)
## Update the path to the files for the local system
dirname(xdata) <- c(rep(system.file("cdf", "KO", package = "faahKO"), 4),</pre>
                    rep(system.file("cdf", "WT", package = "faahKO"), 4))
## Subset to the first 3 files.
xdata <- filterFile(xdata, 1:3, keepFeatures = TRUE)</pre>
## Define features for which to extract EICs
fts <- c("FT097", "FT163", "FT165")
chrs <- featureChromatograms(xdata, features = fts)</pre>
plotChromatogramsOverlay(chrs)
## plot the overlay of EICs in the first sample
plotChromatogramsOverlay(chrs[, 1])
## Define a different color for each feature (row in chrs). By default, also
## all chromatographic peaks of a feature is labeled in the same color.
plotChromatogramsOverlay(chrs[, 1],
    col = c("#ff000040", "#00ff0040", "#0000ff40"))
## Alternatively, we can define a color for each individual chromatographic
## peak and provide this with the `peakBg` and `peakCol` parameters.
chromPeaks(chrs[, 1])
## Use a color for each of the two identified peaks in that sample
plotChromatogramsOverlay(chrs[, 1],
    col = c("#ff000040", "#00ff0040", "#0000ff40"),
    peakBg = c("#ffff0020", "#00ffff20"))
## Plotting the data in all samples.
plotChromatogramsOverlay(chrs,
    col = c("#ff000040", "#00ff0040", "#0000ff40"))
```

```
## Creating a "stacked" EIC plot: the EICs are placed along the y-axis
## relative to their m/z value. With `stacked = 1` the y-axis is split in
## half, the lower half being used for the stacking of the EICs, the upper
## half being used for the *original* intensity axis.
res <- plotChromatogramsOverlay(chrs[, 1], stacked = 1,
    col = c("#ff000040", "#00ff0040", "#0000ff40"))
## add horizontal lines for the m/z values of each EIC
abline(h = res[[1]], col = "grey", lty = 2)
## Note that this type of visualization is different than the conventional
## plot function for chromatographic data, which will draw the EICs for
## multiple samples into the same plot
plot(chrs)
## Converting the object to a MChromatograms without detected peaks
chrs <- as(chrs, "MChromatograms")</pre>
plotChromatogramsOverlay(chrs,
   col = c("#ff000040", "#00ff0040", "#0000ff40"))
```

plotChromPeakDensity,XCMSnExp-method *Plot chromatographic peak density along the retention time axis*

Description

Plot the density of chromatographic peaks along the retention time axis and indicate which peaks would be (or were) grouped into the same feature based using the *peak density* correspondence method. Settings for the *peak density* method can be passed with an PeakDensityParam object to parameter param. If the object contains correspondence results and the correspondence was performed with the *peak groups* method, the results from that correspondence can be visualized setting simulate = FALSE.

Usage

```
## S4 method for signature 'XCMSnExp'
plotChromPeakDensity(
   object,
   mz,
   rt,
   param,
   simulate = TRUE,
   col = "#0000080",
   xlab = "retention time",
   ylab = "sample",
   xlim = range(rt),
   main = NULL,
```

plotChromPeakDensity,XCMSnExp-method

```
type = c("any", "within", "apex_within"),
...
```

Arguments

)

object	A XCMSnExp object with identified chromatographic peaks.
mz	numeric(2) defining an mz range for which the peak density should be plotted.
rt	numeric(2) defining an optional rt range for which the peak density should be plotted. Defaults to the absolute retention time range of object.
param	PeakDensityParam from which parameters for the <i>peak density</i> correspondence algorithm can be extracted. If not provided and if object contains feature definitions with the correspondence/ peak grouping being performed by the <i>peak density</i> method, the corresponding parameter class stored in object is used.
simulate	logical(1) defining whether correspondence should be simulated within the specified m/z / rt region or (with simulate = FALSE) whether the results from an already performed correspondence should be shown.
col	Color to be used for the individual samples. Length has to be 1 or equal to the number of samples in object.
xlab	character(1) with the label for the x-axis.
ylab	character(1) with the label for the y-axis.
xlim	numeric(2) representing the limits for the x-axis. Defaults to the range of the rt parameter.
main	character(1) defining the title of the plot. By default (for main = NULL) the mz-range is used.
type	character(1) specifying how peaks are called to be located within the region defined by mz and rt. Can be one of "any", "within", and "apex_within" for all peaks that are even partially overlapping the region, peaks that are completely within the region, and peaks for which the apex is within the region. This parameter is passed to the chromPeaks function. See related documentation for more information and examples.
	Additional parameters to be passed to the plot function. Data point specific parameters such as bg or pch have to be of length 1 or equal to the number of samples.

Details

The plotChromPeakDensity function allows to evaluate different settings for the *peak density* on an mz slice of interest (e.g. containing chromatographic peaks corresponding to a known metabolite). The plot shows the individual peaks that were detected within the specified mz slice at their retention time (x-axis) and sample in which they were detected (y-axis). The density function is plotted as a black line. Parameters for the density function are taken from the param object. Grey rectangles indicate which chromatographic peaks would be grouped into a feature by the peak density correspondence method. Parameters for the algorithm are also taken from param. See groupChromPeaks-density() for more information about the algorithm and its supported settings.

The function is called for its side effect, i.e. to create a plot.

Author(s)

Johannes Rainer

See Also

groupChromPeaks-density() for details on the *peak density* correspondence method and supported settings.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## Plot the chromatographic peak density for a specific mz range to evaluate
## different peak density correspondence settings.
mzr <- c(305.05, 305.15)
plotChromPeakDensity(faahko_sub, mz = mzr, pch = 16,
    param = PeakDensityParam(sampleGroups = rep(1, length(fileNames(faahko_sub)))))</pre>
```

plotChromPeaks General visualizations of peak detection results

Description

plotChromPeaks plots the identified chromatographic peaks from one file into the plane spanned by the retention time and mz dimension (x-axis representing the retention time and y-axis mz). Each chromatographic peak is plotted as a rectangle representing its width in rt and mz dimension.

This plot is supposed to provide some initial overview of the chromatographic peak detection results.

plotChromPeakImage plots the number of detected peaks for each sample along the retention time axis as an *image* plot, i.e. with the number of peaks detected in each bin along the retention time represented with the color of the respective cell.

Usage

```
plotChromPeaks(
    x,
    file = 1,
    xlim = NULL,
```

plotChromPeaks

```
ylim = NULL,
add = FALSE,
border = "#0000060",
col = NA,
xlab = "retention time",
ylab = "mz",
main = NULL,
...
)
plotChromPeakImage(
    x,
    binSize = 30,
    xlim = NULL,
    log = FALSE,
    xlab = "retention time",
```

main = "Chromatographic peak counts",

yaxt = par("yaxt"),

```
...
)
```

Arguments

x	XCMSnExp object.
file	For plotChromPeaks: numeric(1) specifying the index of the file within x for which the plot should be created. Defaults to 1.
xlim	numeric(2) specifying the x-axis limits (retention time dimension). Defaults to NULL in which case the full retention time range of the file is used.
ylim	For plotChromPeaks: numeric(2) specifying the y-axis limits (mz dimension). Defaults to NULL in which case the full mz range of the file is used.
add	For plotChromPeaks: logical(1) whether the plot should be added or created as a new plot.
border	For plotChromPeaks: the color for the rectangles' border.
col	For plotChromPeaks: the color to be used to fill the rectangles.
xlab	character(1) defining the x-axis label.
ylab	For plotChromPeaks: character(1) defining the y-axis label.
main	character(1) defining the plot title. By default (i.e. main = NULL the name of the file will be used as title.
	Additional arguments passed to the plot (for plotChromPeaks) and image (for plotChromPeakImage) functions. Ignored if add = TRUE.
binSize	For plotChromPeakImage: numeric(1) defining the size of the bins along the x-axis (retention time). Defaults to binSize = 30, peaks within each 30 seconds will thus counted and plotted.
log	For plotChromPeakImage: logical(1) whether the peak counts should be log2 transformed before plotting.

yaxt For plotChromPeakImage: character(1) defining whether y-axis labels should be added. To disable the y-axis use yaxt = "n". For any other value of yaxt the axis will be drawn. See par help page for more details.

Details

The width and line type of the rectangles indicating the detected chromatographic peaks for the plotChromPeaks function can be specified using the par function, i.e. with par(1wd = 3) and par(1ty = 2), respectively.

Author(s)

Johannes Rainer

See Also

highlightChromPeaks for the function to highlight detected chromatographic peaks in extracted ion chromatogram plots.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/KO", package = "faahKO")
## plotChromPeakImage: plot an image for the identified peaks per file
plotChromPeakImage(faahko_sub)
## Show all detected chromatographic peaks from the first file
plotChromPeaks(faahko_sub)
## Plot all detected peaks from the second file and restrict the plot to a
## mz-rt slice
plotChromPeaks(faahko_sub, file = 2, xlim = c(3500, 3600), ylim = c(400, 600))</pre>
```

plotEIC-methods *Plot extracted ion chromatograms for specified m/z range*

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.

plotFeatureGroups

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

Author(s)

Ralf Tautenhahn

See Also

rawEIC,xcmsRaw-class

plotFeatureGroups *Plot feature groups in the m/z-retention time space*

Description

plotFeatureGroups visualizes defined feature groups in the m/z by retention time space. Features are indicated by points with features from the same feature group being connected by a line. See featureGroups() for details on and options for feature grouping.

Usage

```
plotFeatureGroups(
    x,
    xlim = numeric(),
    ylim = numeric(),
    xlab = "retention time",
    ylab = "m/z",
    pch = 4,
```

```
col = "#0000060",
type = "o",
main = "Feature groups",
featureGroups = character()
)
```

х	XCMSnExp() object with grouped features (i.e. after calling groupFeatures().
xlim	numeric(2) with the lower and upper limit for the x-axis.
ylim	numeric(2) with the lower and upper limit for the y-axis.
xlab	character(1) with the label for the x-axis.
ylab	character(1) with the label for the y-axis.
pch	the plotting character. Defaults to pch = 4 i.e. plotting features as crosses. See par() for more information.
col	color to be used to draw the features. At present only a single color is supported.
type	plotting type (see par()). Defaults to type = "o" which draws each feature as a point and connecting the features of the same feature group with a line.
main	character(1) with the title of the plot.
featureGroups	optional character of feature group IDs to draw only specified feature group(s). If not provided, all feature groups are drawn.

Author(s)

Johannes Rainer

plotMsData	DEPRECATED: Create a plot that combines a XIC and a mz/rt 2D
	plot for one sample

Description

UPDATE: please use plot(x,type = "XIC") from the MSnbase package instead. See examples below.

The plotMsData creates a plot that combines an (base peak) extracted ion chromatogram on top (rt against intensity) and a plot of rt against m/z values at the bottom.

Usage

```
plotMsData(
    x,
    main = "",
    cex = 1,
    mfrow = c(2, 1),
    grid.color = "lightgrey",
    colramp = colorRampPalette(rev(brewer.pal(9, "YlGnBu")))
)
```

X	data.frame such as returned by the extractMsData() function. Only a single data.frame is supported.
main	character(1) specifying the title.
cex	numeric(1) defining the size of points. Passed directly to the plot function.
mfrow	<pre>numeric(2) defining the plot layout. This will be passed directly to par(mfrow = mfrow). See par for more information. Setting mfrow = NULL avoids calling par(mfrow = mfrow) hence allowing to pre-define the plot layout.</pre>
grid.color	a color definition for the grid line (or NA to skip creating them).
colramp	a <i>color ramp palette</i> to be used to color the data points based on their intensity. See argument col.regions in lattice::level.colors documentation.

Author(s)

Johannes Rainer

Examples

```
## Read two files from the faahKO package
library(faahKO)
library(magrittr)
cdfs <- dir(system.file("cdf", package = "faahKO"), full.names = TRUE,
    recursive = TRUE)[1:2]
raw_data <- readMSData(cdfs, mode = "onDisk")
## Subset the object to a rt and mz range and plot the data.
raw_data %>%
    filterRt(rt = c(2700, 2900)) %>%
    filterMz(mz = c(334.9, 335.1)) %>%
    plot(type = "XIC")
```

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

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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 outliers

plotRaw-methods

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

List with four matrices, each of dimension features * samples: "mz": median mz deviation for each sample "mzdev": median mz deviation for each sample "rt": median RT deviation for each sample "rtdev": median RT deviation for each sample

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 Scatterplot of raw data points

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

See Also

xcmsRaw-class

DIOLFL-methods Piot retention time deviation brom	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.

Arguments

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

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

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 $\mbox{m}/\mbox{z},$ 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 Plot total ion count

Description

Plot chromatogram of total ion count. Optionally allow identification of target peaks and viewing/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

ProcessHistory-class Tracking data processing

Description

Objects of the type ProcessHistory allow to keep track of any data processing step in an metabolomics experiment. They are created by the data processing methods, such as findChromPeaks and added to the corresponding results objects. Thus, usually, users don't need to create them.

The XProcessHistory extends the ProcessHistory by adding a slot param that allows to store the actual parameter class of the processing step.

processParam, processParam<-: get or set the parameter class from an XProcessHistory object.

msLevel: returns the MS level on which a certain analysis has been performed, or NA if not defined.

The processType method returns a character specifying the processing step type.

The processDate extracts the start date of the processing step.

The processInfo extracts optional additional information on the processing step.

The fileIndex extracts the indices of the files on which the processing step was applied.

Usage

```
## S4 method for signature 'ProcessHistory'
show(object)
## S4 method for signature 'XProcessHistory'
show(object)
## S4 method for signature 'XProcessHistory'
processParam(object)
## S4 method for signature 'ProcessHistory'
msLevel(object)
## S4 method for signature 'ProcessHistory'
processDate(object)
## S4 method for signature 'ProcessHistory'
processInfo(object)
## S4 method for signature 'ProcessHistory'
```

Arguments

fileIndex(object)

object A ProcessHistory or XProcessHistory object.

Value

For processParam: a parameter object extending the Param class.

The processType method returns a character string with the processing step type.

The processDate method returns a character string with the time stamp of the processing step start.

The processInfo method returns a character string with optional additional informations.

The fileIndex method returns a integer vector with the index of the files/samples on which the processing step was applied.

Slots

- type character(1): string defining the type of the processing step. This string has to match predefined values. Use processHistoryTypes to list them.
- date character(1): date time stamp when the processing step was started.
- info character(1): optional additional information.
- fileIndex integer of length 1 or > 1 to specify on which samples of the object the processing was performed.

error (ANY): used to store eventual calculation errors.

param (Param): an object of type Param (e.g. CentWaveParam) specifying the settings of the processing step.

msLevel: integer definining the MS level(s) on which the analysis was performed.

Author(s)

Johannes Rainer

profMat-xcmsSet The profile matrix

Description

The *profile* matrix is an n x m matrix, n (rows) representing equally spaced m/z values (bins) and m (columns) the retention time of the corresponding scans. Each cell contains the maximum intensity measured for the specific scan and m/z values falling within the m/z bin.

The profMat method creates a new profile matrix or returns the profile matrix within the object's @env slot, if available. Settings for the profile matrix generation, such as step (the bin size), method or additional settings are extracted from the respective slots of the xcmsRaw object. Alternatively it is possible to specify all of the settings as additional parameters.

Usage

S4 method for signature 'xcmsRaw'
profMat(object, method, step, baselevel, basespace, mzrange.)

Arguments

object	The xcmsRaw object.
method	The profile matrix generation method. Allowed are "bin", "binlin", "binlinbase" and "intlin". See details section for more information.
step	numeric(1) representing the m/z bin size.
baselevel	numeric(1) representing the base value to which empty elements (i.e. m/z bins without a measured intensity) should be set. Only considered if method = "binlinbase". See baseValue parameter of imputeLinInterpol for more details.
basespace	<pre>numeric(1) representing the m/z length after which the signal will drop to the base level. Linear interpolation will be used between consecutive data points falling within 2 * basespace to each other. Only considered if method = "binlinbase". If not specified, it defaults to 0.075. Internally this parameter is translated into the distance parameter of the imputeLinInterpol function by distance = floor(basespace / step). See distance parameter of imputeLinInterpol for more details.</pre>
mzrange.	Optional numeric(2) manually specifying the mz value range to be used for bin- nind. If not provided, the whole mz value range is used.

Details

Profile matrix generation methods:

- **bin** The default profile matrix generation method that does a simple binning, i.e. aggregating of intensity values falling within an m/z bin.
- **binlin** Binning followed by linear interpolation to impute missing values. The value for m/z bins without a measured intensity are inferred by a linear interpolation between neighboring bins with a measured intensity.
- **binlinbase** Binning followed by a linear interpolation to impute values for empty elements (m/z bins) within a user-definable proximity to non-empty elements while stetting the element's value to the baselevel otherwise. See impute = "linbase" parameter of imputeLinInterpol for more details.
- intlin Set the elements' values to the integral of the linearly interpolated data from plus to minus half the step size.

Value

profMat returns the profile matrix (rows representing scans, columns equally spaced m/z values).

Note

From xcms version 1.51.1 on only the profMat method should be used to extract the profile matrix instead of the previously default way to access it directly *via* object@env\$profile.

Author(s)

Johannes Rainer

See Also

xcmsRaw, binYonX and imputeLinInterpol for the employed binning and missing value imputation methods, respectively. profMat, XCMSnExp-method for the method on XCMSnExp objects.

Examples

```
file <- system.file('cdf/KO/ko15.CDF', package = "faahKO")
## Load the data without generating the profile matrix (profstep = 0)
xraw <- xcmsRaw(file, profstep = 0)
## Extract the profile matrix
profmat <- profMat(xraw, step = 0.3)
dim(profmat)
## If not otherwise specified, the settings from the xraw object are used:
profinfo(xraw)
## To extract a profile matrix with linear interpolation use
profmat <- profMat(xraw, step = 0.3, method = "binlin")
## Alternatively, the profMethod of the xraw objects could be changed
profMethod(xraw) <- "binlin"
profmat_2 <- profMat(xraw, step = 0.3)
all.equal(profmat, profmat_2)</pre>
```

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 Specif

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

profStep-methods

Methods

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

End(Not run)

```
quantify,XCMSnExp-method
```

Accessing mz-rt feature data values

Description

featureValues, XCMSnExp: extract a matrix for feature values with rows representing features and columns samples. Parameter value allows to define which column from the chromPeaks matrix should be returned. Multiple chromatographic peaks from the same sample can be assigned to a feature. Parameter method allows to specify the method to be used in such cases to chose from which of the peaks the value should be returned. Parameter 'msLevel' allows to choose a specific MS level for which feature values should be returned (given that features have been defined for that MS level).

quantify,XCMSnExp: return the preprocessing results as an SummarizedExperiment object containing the feature abundances as assay matrix, the feature definitions (returned by featureDefinitions) as rowData and the phenotype information as colData. This is an ideal container for further processing of the data. Internally, the featureValues method is used to extract the feature abundances, parameters for that method can be passed to quantify with

Usage

```
## S4 method for signature 'XCMSnExp'
quantify(object, ...)
## S4 method for signature 'XCMSnExp'
featureValues(
    object,
    method = c("medret", "maxint", "sum"),
    value = "into",
    intensity = "into",
    filled = TRUE,
    missing = NA,
    msLevel = integer()
)
```

Arguments

object	A XCMSnExp object providing the feature definitions.
	For quantify: additional parameters to be passed on to the featureValues method.
method	character specifying the method to resolve multi-peak mappings within the same sample, i.e. to define the <i>representative</i> peak for a feature in samples where more than one peak was assigned to the feature. If "medret": select the peak closest to the median retention time of the feature. If "maxint": select the peak yielding the largest signal. If "sum": sum the values (only if value is "into" or "maxo".

value	character specifying the name of the column in chromPeaks(object) that should be returned. Defaults to "into" in which case the integrated peak area is returned. To get the index of the peak in the chromPeaks(object) matrix use "index".
intensity	character specifying the name of the column in the chromPeaks(objects) matrix containing the intensity value of the peak that should be used for the conflict resolution if method = "maxint".
filled	logical(1) specifying whether values for filled-in peaks should be returned or not. If filled = FALSE, an NA is returned in the matrix for the respective peak. See fillChromPeaks for details on peak filling.
missing	how missing values should be reported. Allowed values are NA (the default), a numeric or missing = "rowmin_half". The latter replaces any NA with half of the row's minimal (non-missing) value.
msLevel	for 'featureValues': 'integer' defining the MS level(s) for which feature values should be returned. By default, values for features defined for all MS levels are returned.

Value

For featureValues: a matrix with feature values, columns representing samples, rows features. The order of the features matches the order found in the featureDefinitions(object) DataFrame. The rownames of the matrix are the same than those of the featureDefinitions DataFrame. NA is reported for features without corresponding chromatographic peak in the respective sample(s).

For quantify: a SummarizedExperiment representing the preprocessing results.

Note

This method is equivalent to the groupval for xcmsSet objects. Note that missing = 0 should be used to get the same behaviour as groupval, i.e. report missing values as 0 after a call to fillPeaks.

Author(s)

Johannes Rainer

See Also

XCMSnExp for information on the data object.

featureDefinitions to extract the DataFrame with the feature definitions.

featureChromatograms to extract ion chromatograms for each feature.

hasFeatures to evaluate whether the XCMSnExp provides feature definitions.

groupval for the equivalent method on xcmsSet objects.

rawEIC-methods

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 (i.e. values binned along the M/Z dimension).

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

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

Author(s)

Michael Lawrence

See Also

plotRaw for plotting the raw intensities

reconstructChromPeakSpectra

Data independent acquisition (DIA): reconstruct MS2 spectra

Description

Reconstructs MS2 spectra for each MS1 chromatographic peak (if possible) for data independent acquisition (DIA) data (such as SWATH). See the *LC-MS/MS analysis* vignette for more details and examples.

Usage

```
reconstructChromPeakSpectra(
   object,
   expandRt = 0,
   diffRt = 2,
   minCor = 0.8,
   intensity = "maxo",
   peakId = rownames(chromPeaks(object, msLevel = 1L)),
   BPPARAM = bpparam(),
   return.type = c("MSpectra", "Spectra")
)
```

Arguments

object	XCMSnExp with identified chromatographic peaks.
expandRt	numeric(1) allowing to expand the retention time range for extracted ion chro- matograms by a constant value (for the peak shape correlation). Defaults to expandRt = 0 hence correlates only the signal included in the identified chro- matographic peaks.
diffRt	numeric(1) defining the maximal allowed difference between the retention time of the chromatographic peak (apex) and the retention times of MS2 chromatographic peaks (apex) to consider them as representing candidate fragments of the original ion.
minCor	numeric(1) defining the minimal required correlation coefficient for MS2 chro- matographic peaks to be considered for MS2 spectrum reconstruction.
intensity	character(1) defining the column in the chromPeaks matrix that should be used for the intensities of the reconstructed spectra's peaks. The same value from the MS1 chromatographic peaks will be used as precursorIntensity of the resulting spectra.
peakId	optional character vector with peak IDs (i.e. rownames of chromPeaks) of MS1 peaks for which MS2 spectra should be reconstructed. By default they are reconstructed for all MS1 chromatographic peaks.
BPPARAM	parallel processing setup. See <pre>bpparam()</pre> for more information.
return.type	<pre>character(1) defining the type of the returned object. Can be either return.type = "MSpectra" (the default) to return a MSnbase::MSpectra object or return.type = "Spectra" for the newer Spectra::Spectra object.</pre>

Details

In detail, the function performs for each MS1 chromatographic peak:

- Identify all MS2 chromatographic peaks from the isolation window containing the m/z of the ion (i.e. the MS1 chromatographic peak) with approximately the same retention time than the MS1 peak (accepted rt shift can be specified with the diffRt parameter).
- Correlate the peak shapes of the candidate MS2 chromatographic peaks with the peak shape of the MS1 peak retaining only MS2 chromatographic peaks for which the correlation is > min-Cor.

removeIntensity,Chromatogram-method

 Reconstruct the MS2 spectrum using the m/z of all above selected MS2 chromatographic peaks and their intensity (either "maxo" or "into"). Each MS2 chromatographic peak selected for an MS1 peak will thus represent one mass peak in the reconstructed spectrum.

The resulting MSpectra object provides also the peak IDs of the MS2 chromatographic peaks for each spectrum as well as their correlation value.

Value

Depending on return.type:

- MSpectra() with the reconstructed MS2 spectra for all MS1 peaks in object. Contains empty Spectrum2 objects for MS1 peaks for which reconstruction was not possible (either no MS2 signal was recorded or the correlation of the MS2 chromatographic peaks with the MS1 chromatographic peak was below threshold minCor. MSpectra metadata columns "ms2_peak_id" and "ms2_peak_cor" (of type CharacterList() and NumericList() with length equal to the number of peaks per reconstructed MS2 spectrum) providing the IDs and the correlation of the MS2 chromatographic peaks from which the MS2 spectrum was reconstructed. As retention time the median retention times of all MS2 chromatographic peaks used for the spectrum reconstruction is reported. The MS1 chromatographic peak intensity is reported as the reconstructed spectrum's precursorIntensity value (see parameter intensity above).
- Spectra object (defined in the Spectra package). The same content and information than above.

Author(s)

Johannes Rainer, Michael Witting

See Also

findChromPeaksIsolationWindow() for the function to perform MS2 peak detection in DIA isolation windows and for examples.

Description

removeIntensities allows to remove intensities from chromatographic data matching certain conditions (depending on parameter which). The intensities are actually not *removed* but replaced with NA_real_. To actually **remove** the intensities (and the associated retention times) use clean() afterwards.

Parameter which allows to specify which intensities should be replaced by NA_real_. By default (which = "below_threshod" intensities below threshold are removed. If x is a XChromatogram or XChromatograms object (and hence provides also chromatographic peak definitions within the object) which = "outside_chromPeak" can be selected which removes any intensity which is outside the boundaries of identified chromatographic peak(s) in the chromatographic data.

Note that filterIntensity() might be a better approach to subset/filter chromatographic data.

Usage

```
## S4 method for signature 'Chromatogram'
removeIntensity(object, which = "below_threshold", threshold = 0)
## S4 method for signature 'MChromatograms'
removeIntensity(object, which = "below_threshold", threshold = 0)
## S4 method for signature 'XChromatogram'
removeIntensity(
    object,
    which = c("below_threshold", "outside_chromPeak"),
    threshold = 0
)
```

Arguments

object	an object representing chromatographic data. Can be a Chromatogram(), MChromatograms(), XChromatogram() or XChromatograms() object.
which	character(1) defining the condition to remove intensities. See description for details and options.
threshold	<pre>numeric(1) defining the threshold below which intensities are removed (if which = "below_threshold").</pre>

Value

the input object with matching intensities being replaced by NA.

Author(s)

Johannes Rainer

Examples

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.

retcor.obiwarp

Arguments

object	<pre>xcmsSet-class object</pre>
method	Method to use for retention time correction. 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 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

Arguments

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.

Arguments

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
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"),span
= .2,family = c("gaussian","symmetric"),plottype = c("none","deviation","mdevden"),col
= NULL,ty = NULL)
```

See Also

xcmsSet-class, loess retcor.obiwarp

retexp

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

rla	Calculate relative log abundances rla calculates the relative log
	abundances (RLA, see reference) on a numeric vector.

Description

Calculate relative log abundances

rla calculates the relative log abundances (RLA, see reference) on a numeric vector. rowRla calculates row-wise RLAs.

Usage

rla(x, group, log.transform = TRUE)

rowRla(x, group, log.transform = TRUE)

Arguments

x	numeric (for rla) or matrix (for rowRla) with the abundances (in natural scale) on which the RLA should be calculated.
group	factor, numeric or character with the same length than x that groups values in x. If omitted all values are considered to be from the same group.
log.transform	<pre>logical(1) whether x should be log2 transformed. Set to log.transform = FALSE if x is already in log scale.</pre>

Details

The RLA is defines as the (log) abundance of an analyte relative to the median across all abundances of the same group.

Value

numeric of the same length than x (for rla) or matrix with the same dimensions than x (for rowRla).

Author(s)

Johannes Rainer

References

De Livera AM, Dias DA, De Souza D, Rupasinghe T, Pyke J, Tull D, Roessner U, McConville M, Speed TP. Normalizing and integrating metabolomics data. *Anal Chem* 2012 Dec 18;84(24):10768-76.

Examples

x <- c(3, 4, 5, 1, 2, 3, 7, 8, 9)
grp <- c(1, 1, 1, 2, 2, 2, 3, 3, 3)
rla(x, grp)</pre>

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

showError,xcmsSet-method

Extract processing errors

Description

If peak detection is performed with findPeaks setting argument stopOnError = FALSE eventual errors during the process do not cause to stop the processing but are recorded inside of the resulting xcmsSet object. These errors can be accessed with the showError method.

Usage

S4 method for signature 'xcmsSet'
showError(object, message. = TRUE, ...)

Arguments

object	An xcmsSet object.
message.	Logical indicating whether only the error message, or the error itself should be returned.
	Additional arguments.

Value

A list of error messages (if message. = TRUE) or errors or an empty list if no errors are present.

Author(s)

Johannes Rainer

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="meanMZmatch" 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\$specDist.methods. If the nickname of a method is called "meanMZmatch", the help page for that specific method can be accessed with ?specDist.meanMZmatch.

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

Description

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

Usage

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

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

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,matchdist=1,matrix'')
```

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,symmetr
)
```

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.

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

х	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 Divide an xcmsSet object

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

х	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 $h \exp(-(x-mu)^2/(2 \sin ma^2))$, 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.

Arguments

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

stitch-methods

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)

updateObject,xcmsSet-method

Update an xcmsSet object

Description

This method updates an *old* xcmsSet object to the latest definition.

Usage

```
## S4 method for signature 'xcmsSet'
updateObject(object, ..., verbose = FALSE)
```

Arguments

object	The xcmsSet object to update.
	Optional additional arguments. Currently ignored.
verbose	Currently ignored.

Value

An updated xcmsSet containing all data from the input object.

Author(s)

Johannes Rainer

useOriginalCode Enable usage of old xcms code

Description

This function allows to enable the usage of old, partially deprecated code from xcms by setting a corresponding global option. See details for functions affected.

Usage

useOriginalCode(x)

Arguments

Х

logical(1) to specify whether or not original old code should be used in corresponding functions. If not provided the function simply returns the value of the global option.

Details

The functions/methods that are affected by this option are:

- do_findChromPeaks_matchedFilter: use the original code that iteratively creates a subset of the binned (profile) matrix. This is helpful for computers with limited memory or matched-Filter settings with a very small bin size.
- getPeaks

Value

logical(1) indicating whether old code is being used.

Note

For parallel processing using the SOCKS method (e.g. by SnowParam() on Windows computers) this option might not be passed to the individual R processes performing the calculations. In such cases it is suggested to specify the option manually and system-wide by adding the line options(XCMSuseOriginalCode = TRUE) in a file called *.Rprofile* in the folder in which new R processes are started (usually the user's home directory; to ensure that the option is correctly read add a new line to the file too). See also Startup from the base R documentation on how to specify system-wide options for R.

Usage of old code is strongly dicouraged. This function is thought to be used mainly in the transition phase from xcms to xcms version 3.

Author(s)

Johannes Rainer

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.

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,

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,

Description

writeMSData exports mass spectrometry data in mzML or mzXML format. If adjusted retention times are present, these are used as retention time of the exported spectra.

Usage

```
## S4 method for signature 'XCMSnExp,character'
writeMSData(
    object,
    file,
    outformat = c("mzml", "mzxml"),
    copy = FALSE,
    software_processing = NULL,
    ...
)
```

Arguments

object	XCMSnExp object with the mass spectrometry data.
file	character with the file name(s). The length of this parameter has to match the number of files/samples of object.
outformat	character(1) defining the format of the output files (either "mzml" or "mzxml").
сору	logical(1) if metadata (data processing, software used, original file names etc) should be copied from the original files.

writeMzTab

software_processing		
	optionally provide specific data processing steps. See documentation of the software_processing parameter of mzR::writeMSData().	
	Additional parameters to pass down to the writeMSData() function in the MSnbase package, such as outformat to specify the output format ("mzml" or "mzxml") or copy to specify whether general information from the original MS data files (such as data processing, software etc) should be copied to the new files.	

Author(s)

Johannes Rainer

See Also

writeMSData() function in the MSnbase package.

writeMzTab Save a grouped xcmsSet object in mzTab-1.1 format file	
---	--

Description

Write the grouped xcmsSet to an mzTab file.

Arguments

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

Details

The mzTab file format for MS-based metabolomics (and proteomics) is a lightweight supplement to the existing standard XML-based file formats (mzML, mzIdentML, mzQuantML), providing a comprehensive summary, similar in concept to the supplemental material of a scientific publication. mzTab files from xcms contain small molecule sections together with experimental metadata and basic quantitative information. The format is intended to store a simple summary of the final results.

Value

None.

Usage

object = "xcmsSet" writeMzTab(object,filename)

See Also

xcmsSet-class, xcmsSet,

Examples

XChromatograms Containers for chromatographic and peak detection data

Description

The XChromatogram object allows to store chromatographic data (e.g. an extracted ion chromatogram) along with identified chromatographic peaks within that data. The object inherits all functions from the Chromatogram() object in the MSnbase package.

Multiple XChromatogram objects can be stored in a XChromatograms object. This class extends MChromatograms() from the MSnbase package and allows thus to arrange chromatograms in a matrix-like structure, columns representing samples and rows m/z-retention time ranges.

All functions are described (grouped into topic-related sections) after the Arguments section.

Usage

XChromatograms(data, phenoData, featureData, chromPeaks, chromPeakData, ...)

```
XChromatogram(
  rtime = numeric(),
  intensity = numeric(),
  mz = c(NA_real_, NA_real_),
  filterMz = c(NA_real_, NA_real_),
  precursorMz = c(NA_real_, NA_real_),
  productMz = c(NA_real_, NA_real_),
  fromFile = integer(),
  aggregationFun = character(),
  msLevel = 1L,
  chromPeaks,
  chromPeakbata
)
## S4 method for signature 'XChromatogram'
show(object)
```

```
## S4 method for signature 'XChromatogram'
chromPeaks(
 object,
  rt = numeric(),
 mz = numeric(),
 ppm = 0,
  type = c("any", "within", "apex_within"),
 msLevel
)
## S4 replacement method for signature 'XChromatogram'
chromPeaks(object) <- value</pre>
## S4 method for signature 'XChromatogram, ANY'
plot(
  х,
  col = "#00000060",
  lty = 1,
  type = "1",
  xlab = "retention time",
 ylab = "intensity",
  main = NULL,
  peakType = c("polygon", "point", "rectangle", "none"),
  peakCol = "#00000060",
  peakBg = "#00000020",
 peakPch = 1,
  . . .
)
## S4 method for signature 'XChromatogram'
filterMz(object, mz, ...)
## S4 method for signature 'XChromatogram'
filterRt(object, rt, ...)
## S4 method for signature 'XChromatogram'
hasChromPeaks(object)
## S4 method for signature 'XChromatogram'
dropFilledChromPeaks(object)
## S4 method for signature 'XChromatogram'
chromPeakData(object)
## S4 replacement method for signature 'XChromatogram'
chromPeakData(object) <- value</pre>
## S4 method for signature 'XChromatogram, MergeNeighboringPeaksParam'
```

```
refineChromPeaks(object, param = MergeNeighboringPeaksParam())
## S4 method for signature 'XChromatogram'
filterChromPeaks(object, method = c("keepTop"), ...)
## S4 method for signature 'XChromatograms'
show(object)
## S4 method for signature 'XChromatograms'
hasChromPeaks(object)
## S4 method for signature 'XChromatograms'
hasFilledChromPeaks(object)
## S4 method for signature 'XChromatograms'
chromPeaks(
 object,
  rt = numeric(),
 mz = numeric(),
 ppm = 0,
  type = c("any", "within", "apex_within"),
 msLevel
)
## S4 method for signature 'XChromatograms'
chromPeakData(object)
## S4 method for signature 'XChromatograms'
filterMz(object, mz, ...)
## S4 method for signature 'XChromatograms'
filterRt(object, rt, ...)
## S4 method for signature 'XChromatograms,ANY'
plot(
  х,
  col = "#0000060".
  lty = 1,
  type = "1",
  xlab = "retention time",
 ylab = "intensity",
 main = NULL,
  peakType = c("polygon", "point", "rectangle", "none"),
  peakCol = "#00000060",
 peakBg = "#00000020",
 peakPch = 1,
  . . .
```

```
)
```

```
## S4 method for signature 'XChromatograms'
processHistory(object, fileIndex, type)
## S4 method for signature 'XChromatograms'
hasFeatures(object, ...)
## S4 method for signature 'XChromatograms'
dropFeatureDefinitions(object, ...)
## S4 method for signature 'XChromatograms,PeakDensityParam'
groupChromPeaks(object, param)
## S4 method for signature 'XChromatograms'
featureDefinitions(
  object,
 mz = numeric(),
  rt = numeric(),
  ppm = 0,
  type = c("any", "within", "apex_within")
)
## S4 method for signature 'XChromatograms, ANY, ANY, ANY'
x[i, j, drop = TRUE]
## S4 method for signature 'XChromatograms'
featureValues(
  object,
 method = c("medret", "maxint", "sum"),
  value = "into",
  intensity = "into",
 missing = NA,
  . . .
)
## S4 method for signature 'XChromatograms'
plotChromPeakDensity(
  object,
  param,
  col = "#00000060",
  xlab = "retention time",
 main = NULL,
  peakType = c("polygon", "point", "rectangle", "none"),
  peakCol = "#00000060",
  peakBg = "#00000020",
  peakPch = 1,
  simulate = TRUE,
  . . .
```

```
## S4 method for signature 'XChromatograms'
dropFilledChromPeaks(object)
```

S4 method for signature 'XChromatograms,MergeNeighboringPeaksParam'
refineChromPeaks(object, param = MergeNeighboringPeaksParam())

S4 method for signature 'XChromatograms'
filterChromPeaks(object, method = c("keepTop"), ...)

Arguments

data	For XChromatograms: list of Chromatogram or XChromatogram objects.
phenoData	For XChromatograms: either a data.frame, AnnotatedDataFrame or NAnnotatedDataFrame describing the phenotypical information of the samples.
featureData	For XChromatograms: either a data.frame or AnnotatedDataFrame with ad- ditional information for each row of chromatograms.
chromPeaks	For XChromatogram: matrix with required columns "rt", "rtmin", "rtmax", "into", "maxo" and "sn". For XChromatograms: list, same length than data, with the chromatographic peaks for each chromatogram. Each element has to be a matrix, the ordering has to match the order of the chromatograms in data.
chromPeakData	For XChromatogram: DataFrame with optional additional annotations for each chromatographic peak. The number of rows has to match the number of chromatographic peaks.
	For filterChromPeaks: additional parameters defining how to filter chromato- graphic peaks. See function description below for details.
rtime	For XChromatogram: numeric with the retention times (length has to be equal to the length of intensity).
intensity	For XChromatogram: numeric with the intensity values (length has to be equal to the length of rtime).
	For `featureValues`: `character(1)` specifying the name of the column in `chromPeaks(object)` containing the intensity value of the peak that should be used for the `method = "maxint"` conflict resolution if.
mz	For XChromatogram: numeric(2) representing the m/z value range (min, max) on which the chromatogram was created. This is supposed to contain the <i>real</i> range of m/z values in contrast to the filterMz below. For chromPeaks and featureDefinitions: numeric(2) defining the m/z range for which chro- matographic peaks or features should be returned. For filterMz: numeric(2) defining the m/z range for which chromatographic peaks should be retained.#'
filterMz	For XChromatogram: numeric(2) representing the m/z value range (min, max) that was used to filter the original object on m/z dimension. If not applicable use filterMz = $c(0,0)$.
precursorMz	For XChromatogram: numeric(2) for SRM/MRM transitions. Represents the mz of the precursor ion. See details for more information.

)

productMz	For XChromatogram: numeric(2) for SRM/MRM transitions. Represents the mz of the product. See details for more information.
fromFile	For XChromatogram: integer(1) the index of the file within the OnDiskMSnExp or MSnExp object from which the chromatogram was extracted.
aggregationFun	For XChromatogram: character(1) specifying the function that was used to aggregate intensity values for the same retention time across the m/z range.
msLevel	For XChromatogram: integer with the MS level from which the chromatogram was extracted. For chromPeaks and chromPeakData: extract chromatographic peaks of a certain MS level.
object	An XChromatogram or XChromatograms object.
rt	For chromPeaks and featureDefinitions: numeric(2) defining the retention time range for which chromatographic peaks or features should be returned. For filterRt: numeric(2) defining the retention time range to reduce object to.
ppm	For chromPeaks and featureDefinitions: numeric(1) defining a ppm to expand the provided m/z range.
type	For chromPeaks and featureDefinitions: character(1) defining which peaks or features to return if rt or mz is provided: "any" (default) return all peaks that are even partially overlapping with rt, "within" return peaks that are com- pletely within rt and "apex_within" return peaks which apex is within rt.
	<pre>For `plot`: what type of plot should be used for the chromatogram (such as `"l"` for lines, `"p"` for points etc), see help of [plot()] in the `graphics` package for more details. For `processHistory`: restrict returned processing steps to specific types. Use [processHistoryTypes()] to list all supported values.</pre>
value	For chromPeaks<-: a numeric matrix with required columns "rt", "rtmin", "rtmax", "into" and "maxo".
	For `featureValues`: `character(1)` specifying the name of the column in `chromPeaks(object)` that should be returned or `"index"` (default) to return the index of the peak associated with the feature in each sample. To return the integrated peak area instead of the index use `value = "into"`.
x	For plot: an XChromatogram or XChromatograms object.
col	For plot: the color to be used to draw the chromatogram.
lty	For plot and plotChromPeakDensity: the line type.
xlab	For plot and plotChromPeakDensity: the x axis label.
ylab	For plot: the y axis label.
main	For plot and plotChromPeakDensity: an optional title for the plot.
реакТуре	For plot and plotChromPeakDensity: character(1) defining how (and if) identified chromatographic peak within the chromatogram should be plotted. Options are "polygon" (default): draw the peak borders with the peakCol color and fill the peak area with the peakBg color, "point": indicate the peak's apex with a point, "rectangle": draw a rectangle around the identified peak and "none": don't draw peaks.

peakCol	For plot and plotChromPeakDensity: the foreground color for the peaks. For peakType = "polygon" and peakType = "rectangle" this is the color for the border. Use NA to not use a foreground color. This should either be a single color or a vector of colors with the same length than chromPeaks(x) has rows.
peakBg	For plot and plotChromPeakDensity: the background color for the peaks. For peakType = "polygon" and peakType = "rectangle" the peak are or rectangle will be filled with this color. Use NA to skip. This should be either a single color or a vector of colors with the same length than chromPeaks(x) has rows.
peakPch	For plot and plotChromPeakDensity: the point character to be used for peakType = "point". See plot() in the graphics package for more details.
param	For groupChromPeaks and plotChromPeakDensity: a PeakDensityParam() object with the settings for the <i>peak density</i> correspondence analysis algorithm.
method	For featureValues: character(1) specifying the method to resolve multi- peak mappings within the sample sample, i.e. to select the <i>representative</i> peak for a feature for which more than one peak was assigned in one sample. Options are "medret" (default): select the peak closest to the median retention time of the feature, "maxint": select the peak with the largest signal and "sum": sum the values of all peaks (only if value is "into" or "maxo"). For filterChromPeaks: character(1) defining the method that should be used to filter chromatographic peaks. See help on filterChromPeaks below for details.
fileIndex	For processHistory: optional integer specifying the index of the files/samples for which the ProcessHistory objects should be returned.
i	For [: integer with the row indices to subset the XChromatograms object.
j	For [: integer with the column indices to subset the XChromatograms object.
drop	For [: logical(1) whether the dimensionality should be dropped (if possible). Defaults to drop = TRUE, thus, if length of i and j is 1 a XChromatogram is returned. Note that drop is ignored if length of i or j is larger than 1, thus a XChromatograms is returned.
missing	For featureValues: how missing values should be reported. Allowed values are NA (default), a numeric(1) to replace NAs with that value or missing = "rowmin_half" to replace NAs with half of the row's minimal (non-missing) value.
simulate	For plotChromPeakDensity: logical(1) whether a correspondence analysis should be <i>simulated</i> based on the available data and the provided PeakDensityParam() param argument. See section <i>Correspondence analysis</i> for details.

Value

See help of the individual functions.

Creation of objects

Objects can be created with the contructor function XChromatogram and XChromatograms, respectively. Also, they can be coerced from Chromatogram or MChromatograms() objects using as(object, "XChromatogram") or as(object, "XChromatograms").

XChromatograms

Filtering and subsetting

Besides classical subsetting with [specific filter operations on MChromatograms() and XChromatograms objects are available. See filterColumnsIntensityAbove() for more details.

- [allows to subset a XChromatograms object by row (i) and column (j), with i and j being of type integer. The featureDefinitions will also be subsetted accordingly and the peakidx column updated.
- filterMz filters the chromatographic peaks within an XChromatogram or XChromatograms, if a column "mz" is present in the chromPeaks matrix. This would be the case if the XChromatogram was extracted from an XCMSnExp() object with the chromatogram() function. All chromatographic peaks with their m/z within the m/z range defined by mz will be retained. Also feature definitions (if present) will be subset accordingly. The function returns a filtered XChromatogram or XChromatograms object.
- filterRt filters chromatogram(s) by the provided retention time range. All eventually present chromatographic peaks with their apex within the retention time range specified with rt will be retained. Also feature definitions, if present, will be filtered accordingly. The function returns a filtered XChromatogram or XChromatograms object.

Accessing data

See also help of Chromatogram in the MSnbase package for general information and data access. The methods listed here are specific for XChromatogram and XChromatograms objects.

- chromPeaks, chromPeaks<-: extract or set the matrix with the chromatographic peak definitions. Parameter rt allows to specify a retention time range for which peaks should be returned along with parameter type that defines how *overlapping* is defined (parameter description for details). For XChromatogram objects the function returns a matrix with columns "rt" (retention time of the peak apex), "rtmin" (the lower peak boundary), "rtmax" (the upper peak boundary), "into" (the ingegrated peak signal/area of the peak), "maxo" (the maximum instensity of the peak and "sn" (the signal to noise ratio). Note that, depending on the peak detection algorithm, the matrix may contain additional columns. For XChromatograms objects the matrix contains also columns "row" and "column" specifying in which chromatogram of object the peak was identified. Chromatographic peaks are ordered by row.
- chromPeakData, chromPeakData<-: extract or set the DataFrame() with optional chromatographic peak annotations.
- hasChromPeaks: infer whether a XChromatogram (or XChromatograms) has chromatographic peaks. For XChromatogram: returns a logical(1), for XChromatograms: returns a matrix, same dimensions than object with either TRUE or FALSE if chromatographic peaks are available in the chromatogram at the respective position.
- hasFilledChromPeaks: whether a XChromatogram (or a XChromatogram in a XChromatograms) has filled-in chromatographic peaks. For XChromatogram: returns a logical(1), for XChromatograms: returns a matrix, same dimensions than object with either TRUE or FALSE if chromatographic peaks are available in the chromatogram at the respective position.
- dropFilledChromPeaks: removes filled-in chromatographic peaks. See dropFilledChromPeaks() help for XCMSnExp() objects for more information.
- hasFeatures: for XChromatograms objects only: if correspondence analysis has been performed and m/z-rt feature definitions are present. Returns a logical(1).

- dropFeatureDefinitions: for XChrmomatograms objects only: delete any correspondence analysis results (and related process history).
- featureDefinitions: for XChromatograms objects only. Extract the results from the correspondence analysis (performed with groupChromPeaks). Returns a DataFrame with the properties of the defined m/z-rt features: their m/z and retention time range. Columns peakidx and row contain the index of the chromatographic peaks in the chromPeaks matrix associated with the feature and the row in the XChromatograms object in which the feature was defined. Similar to the chromPeaks method it is possible to filter the returned feature matrix with the mz, rt and ppm parameters.
- featureValues: for XChromatograms objects only. Extract the abundance estimates for the individuals features. Note that by default (with parameter value = "index" a matrix of indices of the peaks in the chromPeaks matrix associated to the feature is returned. To extract the integrated peak area use value = "into". The function returns a matrix with one row per feature (in featureDefinitions) and each column being a sample (i.e. column of object). For features without a peak associated in a certain sample NA is returned. This can be changed with the missing argument of the function.
- filterChromPeaks: *filters* chromatographic peaks in object depending on parameter method and method-specific parameters passed as additional arguments with Available methods are:
 - method = "keepTop": keep top n (default n = 1L) peaks in each chromatogram ordered by column order (defaults to order = "maxo"). Parameter decreasing (default decreasing = TRUE) can be used to order peaks in descending (decreasing = TRUE) or ascending (decreasing = FALSE) order to keep the top n peaks with largest or smallest values, respectively.
- processHistory: returns a list of ProcessHistory objects representing the individual performed processing steps. Optional parameters type and fileIndex allow to further specify which processing steps to return.

Plotting and visualizing

- plot draws the chromatogram and highlights in addition any chromatographic peaks present in the XChromatogram or XChromatograms (unless peakType = "none" was specified). To draw peaks in different colors a vector of color definitions with length equal to nrow(chromPeaks(x)) has to be submitted with peakCol and/or peakBg defining one color for each peak (in the order as peaks are in chromPeaks(x)). For base peak chromatograms or total ion chromatograms it might be better to set peakType = "none" to avoid generating busy plots.
- plotChromPeakDensity: visualize *peak density*-based correspondence analysis results. See section *Correspondence analysis* for more details.

Chromatographic peak detection

See findChromPeaks-Chromatogram-CentWaveParam for information.

After chromatographic peak detection it is also possible to *refine* identified chromatographic peaks with the refineChromPeaks method (e.g. to reduce peak detection artifacts). Currently, only peak refinement using the *merge neighboring peaks* method is available (see MergeNeighboringPeaksParam() for a detailed description of the approach.

XChromatograms

Correspondence analysis

Identified chromatographic peaks in an XChromatograms object can be grouped into *features* with the groupChromPeaks function. Currently, such a correspondence analysis can be performed with the *peak density* method (see groupChromPeaks for more details) specifying the algorithm settings with a PeakDensityParam() object. A correspondence analysis is performed separately for each row in the XChromatograms object grouping chromatographic peaks across samples (columns).

The analysis results are stored in the returned XChromatograms object and can be accessed with the featureDefinitions method which returns a DataFrame with one row for each feature. Column "row" specifies in which row of the XChromatograms object the feature was identified.

The plotChromPeakDensity method can be used to visualize *peak density* correspondence results, or to *simulate* a peak density correspondence analysis on chromatographic data. The resulting plot consists of two panels, the upper panel showing the chromatographic data as well as the identified chromatographic peaks, the lower panel the distribution of peaks (the peak density) along the retention time axis. This plot shows each peak as a point with it's peak's retention time on the x-axis, and the sample in which it was found on the y-axis. The distribution of peaks along the retention time axis is visualized with a density estimate. Grouped chromatographic peaks are indicated with grey shaded rectangles. Parameter simulate allows to define whether the correspondence analysis should be simulate(simulate=TRUE, based on the available data and the provided PeakDensityParam() parameter class) or not (simulate=FALSE). For the latter it is assumed that a correspondence analysis has been performed with the *peak density* method on the object. See examples below.

Abundance estimates for each feature can be extracted with the featureValues function using parameter value = "into" to extract the integrated peak area for each feature. The result is a matrix, columns being samples and rows features.

Note

Highlighting the peak area(s) in an XChromatogram or XChromatograms object (plot with peakType = "polygon") draws a polygon representing the displayed chromatogram from the peak's minimal retention time to the maximal retention time. If the XChromatograms was extracted from an XCMSnExp() object with the chromatogram() function this might not represent the actual identified peak area if the m/z range that was used to extract the chromatogram was larger than the peak's m/z.

Author(s)

Johannes Rainer

See Also

findChromPeaks-centWave for peak detection on MChromatograms() objects.

Examples

```
## Create an XChromatograms without peak data
xchrs <- XChromatograms(dta)</pre>
## Create an XChromatograms with peaks data
pks <- list(matrix(c(4, 2, 5, 30, 12, NA), nrow = 1,
   dimnames = list(NULL, c("rt", "rtmin", "rtmax", "into", "maxo", "sn"))),
   NULL)
xchrs <- XChromatograms(dta, chromPeaks = pks)</pre>
## Create an XChromatograms from XChromatogram objects
dta <- lapply(dta, as, "XChromatogram")</pre>
chromPeaks(dta[[1]]) <- pks[[1]]</pre>
xchrs <- XChromatograms(dta, nrow = 1)</pre>
hasChromPeaks(xchrs)
## Loading a test data set with identified chromatographic peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahKO")</pre>
## Subset the dataset to the first and third file.
xod_sub <- filterFile(faahko_sub, file = c(1, 3))</pre>
od <- as(xod_sub, "OnDiskMSnExp")</pre>
## Extract chromatograms for a m/z - retention time slice
chrs <- chromatogram(od, mz = 344, rt = c(2500, 3500))
chrs
## ------ ##
##
       Chromatographic peak detection
                                                     ##
## ------ ##
## Perform peak detection using CentWave
xchrs <- findChromPeaks(chrs, param = CentWaveParam())</pre>
xchrs
## Do we have chromatographic peaks?
hasChromPeaks(xchrs)
## Process history
processHistory(xchrs)
## The chromatographic peaks, columns "row" and "column" provide information
## in which sample the peak was identified.
chromPeaks(xchrs)
## Spectifically extract chromatographic peaks for one sample/chromatogram
chromPeaks(xchrs[1, 2])
## Plot the results
```

XChromatograms

plot(xchrs)

```
## Plot the results using a different color for each sample
sample_colors <- c("#ff000040", "#00ff0040", "#0000ff40")</pre>
cols <- sample_colors[chromPeaks(xchrs)[, "column"]]</pre>
plot(xchrs, col = sample_colors, peakBg = cols)
## Indicate the peaks with a rectangle
plot(xchrs, col = sample_colors, peakCol = cols, peakType = "rectangle",
   peakBg = NA)
## ------ ##
    Correspondence analysis
##
                                                     ##
## ------ ##
## Group chromatographic peaks across samples
prm <- PeakDensityParam(sampleGroup = rep(1, 2))</pre>
res <- groupChromPeaks(xchrs, param = prm)</pre>
hasFeatures(res)
featureDefinitions(res)
## Plot the correspondence results. Use simulate = FALSE to show the
## actual results. Grouped chromatographic peaks are indicated with
## grey shaded rectangles.
plotChromPeakDensity(res, simulate = FALSE)
## Simulate a correspondence analysis based on different settings. Larger
## bw will increase the smoothing of the density estimate hence grouping
## chromatographic peaks that are more apart on the retention time axis.
prm <- PeakDensityParam(sampleGroup = rep(1, 3), bw = 60)</pre>
plotChromPeakDensity(res, param = prm)
## Delete the identified feature definitions
res <- dropFeatureDefinitions(res)</pre>
hasFeatures(res)
## Create a XChromatogram object
pks <- matrix(nrow = 1, ncol = 6)</pre>
colnames(pks) <- c("rt", "rtmin", "rtmax", "into", "maxo", "sn")</pre>
pks[, "rtmin"] <- 2
pks[, "rtmax"] <- 9
pks[, "rt"] <- 4
pks[, "maxo"] <- 19
pks[, "into"] <- 93
xchr <- XChromatogram(rtime = 1:10,</pre>
    intensity = c(4, 8, 14, 19, 18, 12, 9, 8, 5, 2),
    chromPeaks = pks)
xchr
## Add arbitrary peak annotations
df <- DataFrame(peak_id = c("a"))</pre>
xchr <- XChromatogram(rtime = 1:10,</pre>
```

```
intensity = c(4, 8, 14, 19, 18, 12, 9, 8, 5, 2),
    chromPeaks = pks, chromPeakData = df)
xchr
chromPeakData(xchr)
## Extract the chromatographic peaks
chromPeaks(xchr)
## Plotting of a single XChromatogram object
## o Don't highlight chromatographic peaks
plot(xchr, peakType = "none")
## o Indicate peaks with a polygon
plot(xchr)
## Add a second peak to the data.
pks <- rbind(chromPeaks(xchr), c(7, 7, 10, NA, 15, NA))</pre>
chromPeaks(xchr) <- pks</pre>
## Plot the peaks in different colors
plot(xchr, peakCol = c("#ff000080", "#0000ff80"),
    peakBg = c("#ff000020", "#0000ff20"))
## Indicate the peaks as rectangles
plot(xchr, peakCol = c("#ff000060", "#0000ff60"), peakBg = NA,
   peakType = "rectangle")
## Filter the XChromatogram by retention time
xchr_sub <- filterRt(xchr, rt = c(4, 6))</pre>
xchr_sub
plot(xchr_sub)
```

xcms-deprecated Deprecated functions in package 'xcms'

Description

These functions are provided for compatibility with older versions of 'xcms' only, and will be defunct at the next release.

Details

The following functions/methods are deprecated.

- profBin, profBinM, profBinLin, profBinLinM, profBinLinBase, profBinLinBaseM have been deprecated and binYonX in combination with imputeLinInterpol should be used instead.
- extractMsData: replaced by as(x, "data.frame").
- plotMsData: replaced by plot(x,type = "XIC").

264

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",...).

Slots

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.

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

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.

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 collisionEnergy for the spectra.

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.

Author(s)

S. Neumann, J. Kutzera

XCMSnExp-class

See Also

xcmsRaw

XCMSnExp-class

Data container storing xcms preprocessing results

Description

The XCMSnExp object is a container for the results of a G/LC-MS data preprocessing that comprises chromatographic peak detection, alignment and correspondence. These results can be accessed with the chromPeaks, adjustedRtime and featureDefinitions functions; see below (after the Usage, Arguments, Value and Slots sections) for more details). Along with the results, the object contains the processing history that allows to track each processing step along with the used settings. This can be extracted with the processHistory method. XCMSnExp objects, by directly extending the OnDiskMSnExp object from the MSnbase package, inherit all of its functionality and allows thus an easy access to the full raw data at any stage of an analysis. To support interaction with packages requiring the *old* objects, XCMSnExp objects can be coerced into xcmsSet objects using the as method (see examples below). All preprocessing results will be passed along to the resulting xcmsSet object.

General functions for XCMSnExp objects are (see further below for specific function to handle chromatographic peak data, alignment and correspondence results):

processHistoryTypes returns the available *types* of process histories. These can be passed with argument type to the processHistory method to extract specific process step(s).

hasFilledChromPeaks: whether filled-in peaks are present or not.

featureArea extracts the m/z - retention time region for each feature. This area is defined by the m/z - retention time regions of all chromatographic peaks associated with a feature. Parameters mzmin, mzmax, rtmin and rtmax allow to define functions how the corresponding value is calculated from the individual values (such as the "rtmin") of all chromatographic peaks of that feature. By default the median "rtmin", "rtmax", "mzmin" and "mzmax" is reported. Parameter features allows to provide feature IDs for which the area should be extracted. By default it is extracted for all features.

profMat: creates a *profile matrix*, which is a n x m matrix, n (rows) representing equally spaced m/z values (bins) and m (columns) the retention time of the corresponding scans. Each cell contains the maximum intensity measured for the specific scan and m/z values. See profMat for more details and description of the various binning methods.

hasAdjustedRtime: whether the object provides adjusted retention times.

hasFeatures: whether the object contains correspondence results (i.e. features).

hasChromPeaks: whether the object contains peak detection results.

hasFilledChromPeaks: whether the object contains any filled-in chromatographic peaks.

adjustedRtime,adjustedRtime<-: extract/set adjusted retention times. adjustedRtime<- should not be called manually, it is called internally by the adjustRtime methods. For XCMSnExp objects, adjustedRtime<- does also apply retention time adjustments to eventually present chromatographic peaks. The bySample parameter allows to specify whether the adjusted retention time should be grouped by sample (file). featureDefinitions, featureDefinitions<-: extract or set the correspondence results, i.e. the mz-rt features (peak groups). Similar to the chromPeaks it is possible to extract features for specified m/z and/or rt ranges. The function supports also the parameter type that allows to specify which features to be returned if any of rt or mz is specified. For details see help of chromPeaks. See also featureSummary for a function to calculate simple feature summaries.

chromPeaks, chromPeaks<-: extract or set the matrix containing the information on identified chromatographic peaks. Rownames of the matrix represent unique IDs of the respective peaks within the experiment. Parameter by Sample allows to specify whether peaks should be returned ungrouped (default bySample = FALSE) or grouped by sample (bySample = TRUE). The chromPeaks<method for XCMSnExp objects removes also all correspondence (peak grouping) and retention time correction (alignment) results. The optional arguments rt, mz, ppm and type allow to extract only chromatographic peaks overlapping the defined retention time and/or m/z ranges. Argument type allows to define how *overlapping* is determined: for type == "any" (the default), all peaks that are even partially overlapping the region are returned (i.e. for which either "mzmin" or "mzmax" of the chromPeaks or featureDefinitions matrix are within the provided m/z range), for type == "within" the full peak has to be within the region (i.e. both "mzmin" and "mzmax" have to be within the m/z range) and for type == "apex_within" the peak's apex position (highest signal of the peak) has to be within the region (i.e. the peak's or features m/z has to be within the m/z range). See description of the return value for details on the returned matrix. Users usually don't have to use the chromPeaks<- method directly as detected chromatographic peaks are added to the object by the findChromPeaks method. Also, chromPeaks<- will replace any existing chromPeakData.

chromPeakData and chromPeakData<- allow to get or set arbitrary chromatographic peak annotations. These are returned or ar returned as a DataFrame. Note that the number of rows and the rownames of the DataFrame have to match those of chromPeaks.

rtime: extracts the retention time for each scan. The bySample parameter allows to return the values grouped by sample/file and adjusted whether adjusted or raw retention times should be returned. By default the method returns adjusted retention times, if they are available (i.e. if retention times were adjusted using the adjustRtime method).

mz: extracts the mz values from each scan of all files within an XCMSnExp object. These values are extracted from the original data files and eventual processing steps are applied *on the fly*. Using the bySample parameter it is possible to switch from the default grouping of mz values by spectrum/scan to a grouping by sample/file.

intensity: extracts the intensity values from each scan of all files within an XCMSnExp object. These values are extracted from the original data files and eventual processing steps are applied *on the fly*. Using the bySample parameter it is possible to switch from the default grouping of intensity values by spectrum/scan to a grouping by sample/file.

spectra: extracts the Spectrum objects containing all data from object. The values are extracted from the original data files and eventual processing steps are applied *on the fly*. By setting bySample = TRUE, the spectra are returned grouped by sample/file. If the XCMSnExp object contains adjusted retention times, these are returned by default in the Spectrum objects (can be overwritten by setting adjusted = FALSE).

processHistory: returns a list of ProcessHistory objects (or objects inheriting from this base class) representing the individual processing steps that have been performed, eventually along with their settings (Param parameter class). Optional arguments fileIndex, type and msLevel allow to restrict to process steps of a certain type or performed on a certain file or MS level.

dropChromPeaks: drops any identified chromatographic peaks and returns the object without that

XCMSnExp-class

information. Note that for XCMSnExp objects the method drops by default also results from a correspondence (peak grouping) analysis. Adjusted retention times are removed if the alignment has been performed *after* peak detection. This can be overruled with keepAdjustedRtime = TRUE.

dropFeatureDefinitions: drops the results from a correspondence (peak grouping) analysis, i.e. the definition of the mz-rt features and returns the object without that information. Note that for XCMSnExp objects the method will also by default drop retention time adjustment results, if these were performed after the last peak grouping (i.e. which base on the results from the peak grouping that are going to be removed). All related process history steps are removed too as well as eventually filled in peaks (by fillChromPeaks). The parameter keepAdjustedRtime can be used to avoid removal of adjusted retention times.

dropAdjustedRtime: drops any retention time adjustment information and returns the object without adjusted retention time. For XCMSnExp objects, this also reverts the retention times reported for the chromatographic peaks in the peak matrix to the original, raw, ones (after chromatographic peak detection). Note that for XCMSnExp objects the method drops also all peak grouping results if these were performed *after* the retention time adjustment. All related process history steps are removed too.

findChromPeaks performs chromatographic peak detection on the provided XCMSnExp objects. For more details see the method for XCMSnExp. Note that by default (with parameter add = FALSE) previous peak detection results are removed. Use add = TRUE to perform a second round of peak detection and add the newly identified peaks to the previous peak detection results. Correspondence results (features) are always removed prior to peak detection. Previous alignment (retention time adjustment) results are kept, i.e. chromatographic peak detection is performed using adjusted retention times if the data was first aligned using e.g. obiwarp (adjustRtime-obiwarp).

dropFilledChromPeaks: drops any filled-in chromatographic peaks (filled in by the fillChromPeaks method) and all related process history steps.

spectrapply applies the provided function to each Spectrum in the object and returns its results. If no function is specified the function simply returns the list of Spectrum objects.

XCMSnExp objects can be combined with the c function. This combines identified chromatographic peaks and the objects' pheno data but discards alignment results or feature definitions.

plot plots the spectrum data (see plot for MSnExp objects in the MSnbase package for more details. For type = "XIC", identified chromatographic peaks will be indicated as rectangles with border color peakCol.

Usage

```
processHistoryTypes()
```

hasFilledChromPeaks(object)

```
featureArea(
   object,
   mzmin = median,
   mzmax = median,
   rtmin = median,
   rtmax = median,
```

```
msLevel = unique(msLevel(object)),
```

```
features = character()
```

```
)
## S4 method for signature 'OnDiskMSnExp'
profMat(
  object,
 method = "bin",
  step = 0.1,
  baselevel = NULL,
  basespace = NULL,
 mzrange. = NULL,
  fileIndex,
  . . .
)
## S4 method for signature 'XCMSnExp'
show(object)
## S4 method for signature 'XCMSnExp'
hasAdjustedRtime(object)
## S4 method for signature 'XCMSnExp'
hasFeatures(object, msLevel = integer())
## S4 method for signature 'XCMSnExp'
hasChromPeaks(object, msLevel = integer())
## S4 method for signature 'XCMSnExp'
hasFilledChromPeaks(object)
## S4 method for signature 'XCMSnExp'
adjustedRtime(object, bySample = FALSE)
## S4 replacement method for signature 'XCMSnExp'
adjustedRtime(object) <- value</pre>
## S4 method for signature 'XCMSnExp'
featureDefinitions(
 object,
 mz = numeric(),
  rt = numeric(),
  ppm = 0,
  type = c("any", "within", "apex_within"),
 msLevel = integer()
)
## S4 replacement method for signature 'XCMSnExp'
```

```
featureDefinitions(object) <- value</pre>
```

272

```
## S4 method for signature 'XCMSnExp'
chromPeaks(
 object,
 bySample = FALSE,
 rt = numeric(),
 mz = numeric(),
 ppm = 0,
 msLevel = integer(),
  type = c("any", "within", "apex_within"),
  isFilledColumn = FALSE
)
## S4 replacement method for signature 'XCMSnExp'
chromPeaks(object) <- value</pre>
## S4 method for signature 'XCMSnExp'
rtime(object, bySample = FALSE, adjusted = hasAdjustedRtime(object))
## S4 method for signature 'XCMSnExp'
mz(object, bySample = FALSE, BPPARAM = bpparam())
## S4 method for signature 'XCMSnExp'
intensity(object, bySample = FALSE, BPPARAM = bpparam())
## S4 method for signature 'XCMSnExp'
spectra(
  object,
 bySample = FALSE,
 adjusted = hasAdjustedRtime(object),
 BPPARAM = bpparam()
)
## S4 method for signature 'XCMSnExp'
processHistory(object, fileIndex, type, msLevel)
## S4 method for signature 'XCMSnExp'
dropChromPeaks(object, keepAdjustedRtime = FALSE)
## S4 method for signature 'XCMSnExp'
dropFeatureDefinitions(object, keepAdjustedRtime = FALSE, dropLastN = -1)
## S4 method for signature 'XCMSnExp'
dropAdjustedRtime(object)
## S4 method for signature 'XCMSnExp'
profMat(
 object,
 method = "bin",
```

```
step = 0.1,
  baselevel = NULL,
 basespace = NULL,
 mzrange. = NULL,
 fileIndex,
  . . .
)
## S4 method for signature 'XCMSnExp,Param'
findChromPeaks(
 object,
  param,
 BPPARAM = bpparam(),
 return.type = "XCMSnExp",
 msLevel = 1L,
  add = FALSE
)
## S4 method for signature 'XCMSnExp'
dropFilledChromPeaks(object)
## S4 method for signature 'XCMSnExp'
spectrapply(object, FUN = NULL, BPPARAM = bpparam(), ...)
## S3 method for class 'XCMSnExp'
c(...)
## S4 method for signature 'XCMSnExp'
chromPeakData(object)
## S4 replacement method for signature 'XCMSnExp'
chromPeakData(object) <- value</pre>
## S4 method for signature 'XCMSnExp,missing'
```

plot(x, y, type = c("spectra", "XIC"), peakCol = "#ff000060", ...)

Arguments

object	For adjustedRtime, featureDefinitions, chromPeaks, hasAdjustedRtime, hasFeatures and hasChromPeaks either a MsFeatureData or a XCMSnExp object, for all other methods a XCMSnExp object.
mzmin	for featureArea: function to be applied to values in the "mzmin" column of all chromatographic peaks of a feature to define the lower m/z value of the feature area. Defaults to median.
mzmax	for featureArea: function same as mzmin but for the "mzmax" column.
rtmin	for featureArea: function same as mzmin but for the "rtmin" column.
rtmax	for featureArea: function same as mzmin but for the "rtmax" column.

274

msLevel	integer specifying the MS level(s) for which identified chromatographic peaks should be returned.
features	for featureArea: IDs of features for which the area should be extracted.
method	The profile matrix generation method. Allowed are "bin", "binlin", "binlinbase" and "intlin". See details section for more information.
step	numeric(1) representing the m/z bin size.
baselevel	numeric(1) representing the base value to which empty elements (i.e. m/z bins without a measured intensity) should be set. Only considered if method = "binlinbase". See baseValue parameter of imputeLinInterpol for more details.
basespace	<pre>numeric(1) representing the m/z length after which the signal will drop to the base level. Linear interpolation will be used between consecutive data points falling within 2 * basespace to each other. Only considered if method = "binlinbase". If not specified, it defaults to 0.075. Internally this parameter is translated into the distance parameter of the imputeLinInterpol function by distance = floor(basespace / step). See distance parameter of imputeLinInterpol for more details.</pre>
mzrange.	Optional numeric(2) manually specifying the mz value range to be used for bin- nind. If not provided, the whole mz value range is used.
fileIndex	For processHistory: optional integer specifying the index of the files/samples for which the ProcessHistory objects should be retrieved.
	Additional parameters.
bySample	logical(1) specifying whether results should be grouped by sample.
value	For adjustedRtime<-: a list (length equal to the number of samples) with numeric vectors representing the adjusted retention times per scan.
	For featureDefinitions<-: a DataFrame with peak grouping information. See return value for the featureDefinitions method for the expected format.
	For chromPeaks<-: a matrix with information on detected peaks. See return value for the chromPeaks method for the expected format.
mz	optional numeric(2) defining the mz range for which chromatographic peaks should be returned.
rt	optional numeric(2) defining the retention time range for which chromato- graphic peaks should be returned.
ppm	optional numeric(1) specifying the ppm by which the mz range should be ex- tended. For a value of ppm = 10, all peaks within mz[1] -ppm / 1e6 and mz[2] + ppm / 1e6 are returned.
type	For processHistory: restrict returned ProcessHistory objects to analysis steps of a certain type. Use the processHistoryTypes to list all supported values. For chromPeaks: character specifying which peaks to return if rt or mz are defined. For type = "any" all chromatographic peaks partially overlap- ping the range defined by mz and/or rt are returned, type = "within" returns only peaks completely within the region and type = "apex_within" peaks for which the peak's apex is within the region.
isFilledColumn	logical(1) whether a column "is_filled" is included in the returned "matrix" providing the information if a peak was filled in. Alternatively, this information would be provided by the chromPeakData data frame.

adjusted	logical(1) whether adjusted or raw (i.e. the original retention times reported in the files) should be returned.		
BPPARAM	Parameter class for parallel processing. See bpparam.		
keepAdjustedRt	keepAdjustedRtime		
	For dropFeatureDefinitions, XCMSnExp: logical(1) defining whether even- tually present retention time adjustment should not be dropped. By default drop- ping feature definitions drops retention time adjustment results too.		
dropLastN	For dropFeatureDefinitions,XCMSnExp: numeric(1) defining the number of peak grouping related process history steps to remove. By default dropLastN = -1, dropping the chromatographic peaks removes all process history steps re- lated to peak grouping. Setting e.g. dropLastN = 1 will only remove the most recent peak grouping related process history step.		
param	A CentWaveParam, MatchedFilterParam, MassifquantParam, MSWParam or CentWavePredIsoParam object with the settings for the chromatographic peak detection algorithm.		
return.type	Character specifying what type of object the method should return. Can be either "XCMSnExp" (default), "list" or "xcmsSet".		
add	For findChromPeaks: if newly identified chromatographic peaks should be added to the peak matrix with the already identified chromatographic peaks. By default (add = FALSE) previous peak detection results will be removed.		
FUN	For spectrapply: a function that should be applied to each spectrum in the object.		
х	For plot: XCMSnExp object.		
у	For plot: not used.		
peakCol	For plot: the color that should be used to indicate identified chromatographic peaks (only in combination with type = "XIC" and if chromatographic peaks are present).		

Value

For profMat: a list with a the profile matrix matrix (or matrices if fileIndex was not specified or if length(fileIndex) > 1). See profile-matrix for general help and information about the profile matrix.

For adjustedRtime: if bySample = FALSE a numeric vector with the adjusted retention for each spectrum of all files/samples within the object. If bySample = TRUE a list (length equal to the number of samples) with adjusted retention times grouped by sample. Returns NULL if no adjusted retention times are present.

For featureDefinitions: a DataFrame with peak grouping information, each row corresponding to one mz-rt feature (grouped peaks within and across samples) and columns "mzmed" (median mz value), "mzmin" (minimal mz value), "mzmax" (maximum mz value), "rtmed" (median retention time), "rtmin" (minimal retention time), "rtmax" (maximal retention time) and "peakidx". Column "peakidx" contains a list with indices of chromatographic peaks (rows) in the matrix returned by the chromPeaks method that belong to that feature group. The method returns NULL if no feature definitions are present. featureDefinitions supports also parameters mz, rt, ppm and type to return only features within certain ranges (see description of chromPeaks for details).

XCMSnExp-class

For chromPeaks: if bySample = FALSE a matrix (each row being a chromatographic peak, rownames representing unique IDs of the peaks) with at least the following columns: "mz" (intensityweighted mean of mz values of the peak across scans/retention times), "mzmin" (minimal mz value), "mzmax" (maximal mz value), "rt" (retention time of the peak apex), "rtmin" (minimal retention time), "rtmax" (maximal retention time), "into" (integrated, original, intensity of the peak), "maxo" (maximum intentity of the peak), "sample" (sample index in which the peak was identified) and Depending on the employed peak detection algorithm and the verboseColumns parameter of it, additional columns might be returned. If parameter isFilledColumn was set to TRUE a column named "is_filled" is also returned. For bySample = TRUE the chromatographic peaks are returned as a list of matrices, each containing the chromatographic peaks of a specific sample. For samples in which no peaks were detected a matrix with 0 rows is returned.

For rtime: if bySample = FALSE a numeric vector with the retention times of each scan, if bySample = TRUE a list of numeric vectors with the retention times per sample.

For mz: if bySample = FALSE a list with the mz values (numeric vectors) of each scan. If bySample = TRUE a list with the mz values per sample.

For intensity: if bySample = FALSE a list with the intensity values (numeric vectors) of each scan. If bySample = TRUE a list with the intensity values per sample.

For spectra: if bySample = FALSE a list with Spectrum objects. If bySample = TRUE the result is grouped by sample, i.e. as a list of lists, each element in the *outer* list being the list of spectra of the specific file.

For processHistory: a list of ProcessHistory objects providing the details of the individual data processing steps that have been performed.

Slots

- .processHistory list with XProcessHistory objects tracking all individual analysis steps that have been performed.
- msFeatureData MsFeatureData class extending environment and containing the results from a chromatographic peak detection (element "chromPeaks"), peak grouping (element "featureDefinitions") and retention time correction (element "adjustedRtime") steps. This object should not be manipulated directly.

Chromatographic peak data

Chromatographic peak data is added to an XCMSnExp object by the findChromPeaks function. Functions to access chromatographic peak data are:

- hasChromPeaks whether chromatographic peak data is available, see below for help of the function.
- chromPeaks access chromatographic peaks (see below for help).
- dropChromPeaks remove chromatographic peaks (see below for help).
- dropFilledChromPeaks remove filled-in peaks (see below for help).
- fillChromPeaks fill-in missing peaks (see respective help page).
- plotChromPeaks plot identified peaks for a file (see respective help page).
- plotChromPeakImage plot distribution of peaks along the retention time axis (see respective help page).

• highlightChromPeaks add chromatographic peaks to an existing plot of a Chromatogram (see respective help page).

Adjusted retention times

Adjusted retention times are stored in an XCMSnExp object besides the original, raw, retention times, allowing to switch between raw and adjusted times. It is also possible to replace the raw retention times with the adjusted ones with the applyAdjustedRtime. The adjusted retention times are added to an XCMSnExp by the adjustRtime function. All functions related to the access of adjusted retention times are:

- hasAdjustedRtime whether adjusted retention times are available (see below for help).
- dropAdjustedRtime remove adjusted retention times (see below for help).
- applyAdjustedRtime replace the raw retention times with the adjusted ones (see respective help page).
- plotAdjustedRtime plot differences between adjusted and raw retention times (see respective help page).

Correspondence results, features

The correspondence analysis (groupChromPeaks) adds the feature definitions to an XCMSnExp object. All functions related to these are listed below:

- hasFeatures whether correspondence results are available (see below for help).
- featureDefinitions access the definitions of the features (see below for help).
- dropFeatureDefinitions remove correspondence results (see below for help).
- featureValues access values for features (see respective help page).
- featureSummary perform a simple summary of the defined features (see respective help page).
- overlappingFeatures identify features that are overlapping or close in the m/z rt space (see respective help page).
- quantify extract feature intensities and put them, along with feature definitions and phenodata information, into a SummarizedExperiment. See help page for details.

Note

The "chromPeaks" element in the msFeatureData slot is equivalent to the @peaks slot of the xcmsSet object, the "featureDefinitions" contains information from the @groups and @groupidx slots from an xcmsSet object.

Author(s)

Johannes Rainer

XCMSnExp-class

See Also

xcmsSet for the old implementation. OnDiskMSnExp, MSnExp and pSet for a complete list of inherited methods.

findChromPeaks for available peak detection methods returning a XCMSnExp object as a result.

groupChromPeaks for available peak grouping methods and featureDefinitions for the method to extract the feature definitions representing the peak grouping results. adjustRtime for retention time adjustment methods.

chromatogram to extract MS data as Chromatogram objects.

as (as(x, "data.frame")) in the MSnbase package for the method to extract MS data as data.frames.

featureSummary to calculate basic feature summaries.

featureChromatograms to extract chromatograms for each feature.

chromPeakSpectra to extract MS2 spectra with the m/z of the precursor ion within the m/z range of a peak and a retention time within its retention time range.

featureSpectra to extract MS2 spectra associated with identified features.

fillChromPeaks for the method to fill-in eventually missing chromatographic peaks for a feature in some samples.

Examples

```
## Load a test data set with detected peaks
data(faahko_sub)
## Update the path to the files for the local system
dirname(faahko_sub) <- system.file("cdf/K0", package = "faahK0")</pre>
```

```
## Disable parallel processing for this example
register(SerialParam())
```

```
## The results from the peak detection are now stored in the XCMSnExp
## object
faahko_sub
```

The detected peaks can be accessed with the chromPeaks method. head(chromPeaks(faahko_sub))

```
## The settings of the chromatographic peak detection can be accessed with
## the processHistory method
processHistory(faahko_sub)
```

```
## Also the parameter class for the peak detection can be accessed
processParam(processHistory(faahko_sub)[[1]])
```

```
## The XCMSnExp inherits all methods from the pSet and OnDiskMSnExp classes
## defined in Bioconductor's MSnbase package. To access the (raw) retention
## time for each spectrum we can use the rtime method. Setting bySample = TRUE
## would cause the retention times to be grouped by sample
head(rtime(faahko_sub))
```

Similarly it is possible to extract the mz values or the intensity values

```
## using the mz and intensity method, respectively, also with the option to
## return the results grouped by sample instead of the default, which is
## grouped by spectrum. Finally, to extract all of the data we can use the
## spectra method which returns Spectrum objects containing all raw data.
## Note that all these methods read the information from the original input
## files and subsequently apply eventual data processing steps to them.
mzs <- mz(faahko_sub, bySample = TRUE)</pre>
length(mzs)
lengths(mzs)
## The full data could also be read using the spectra data, which returns
## a list of Spectrum object containing the mz, intensity and rt values.
## spctr <- spectra(faahko_sub)</pre>
## To get all spectra of the first file we can split them by file
## head(split(spctr, fromFile(faahko_sub))[[1]])
#############
## Filtering
##
## XCMSnExp objects can be filtered by file, retention time, mz values or
## MS level. For some of these filter preprocessing results (mostly
## retention time correction and peak grouping results) will be dropped.
## Below we filter the XCMSnExp object by file to extract the results for
## only the second file.
xod_2 <- filterFile(faahko_sub, file = 2)</pre>
xod_2
## Now the objects contains only the idenfified peaks for the second file
head(chromPeaks(xod_2))
##########
## Coercing to an xcmsSet object
##
## We can also coerce the XCMSnExp object into an xcmsSet object:
xs <- as(faahko_sub, "xcmsSet")</pre>
```

head(peaks(xs))

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", ...).

xcmsRaw

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

Constructor for xcmsRaw objects which reads NetCDF/mzXML files

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. See profile-matrix for details and supported values.
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)

xcmsRaw

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

See profile-matrix for details on profile matrix generation methods and settings.

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>

References

NetCDF file format: https://www.unidata.ucar.edu/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
```

xcmsRaw-class

```
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]]</pre>
    intensity.scan1<-xr@env$intensity[(1+xr@scanindex[1]):xr@scanindex[2]]</pre>
   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)</pre>
   head(scan1)
   plotScan(xr, 1)
## End(Not run)
```

xcmsRaw-class

Class xcmsRaw, a class for handling raw data

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: Numeric representing the acquisition number of the individual scans/spectra. Length of acquisitionNum is equal to the number of spectra/scans in the object and hence equal to the scantime slot. Note however that this information is only available in mzML files.
- 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. The profile matrix should be extracted with the profMat method.

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: list to store additional profile matrix generation settings. Use the profinfo method to extract all profile matrix creation relevant information.
- 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. Note that the scanrange will always be 1 to the number of scans within the xcmsRaw object, which does not necessarily have to match to the scan index in the original mzML file (e.g. if the original data was subsetted). The acquisitionNum information can be used to track the original *position* of each scan in the mzML file.

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

xcmsRaw-class

- **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
- 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</pre>
- 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. See slot description above for more information.
- 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.
- findmzROI signature(object = "xcmsRaw"): internal function to identify regions of interest in the raw data as part of the first step of centWave-based peak detection.

Author(s)

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See Also

xcmsRaw, subset-xcmsRaw for subsetting by spectra.

xcmsSet

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(files = NULL, snames = NULL, sclass = NULL, phenoData = NULL,
    profmethod = "bin", profparam = list(),
    polarity = NULL, lockMassFreq=FALSE,
mslevel=NULL, nSlaves=0, progressCallback=NULL,
    scanrange = NULL, BPPARAM = bpparam(),
    stopOnError = TRUE, ...)
```

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. Supported values are "bin", "binlin", "binlinbase" and "intlin" (for methods profBin, profBinLin, profBinLinBase and profIntLin, respectively). See help on profBin for a complete list of available methods and their supported parameters.	
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	DEPRECATED, use BPPARAM argument instead.	
progressCallback		
	function to be called, when progressInfo changes (useful for GUIs)	
scanrange	scan range to read	
BPPARAM	a BiocParallel parameter object to control how and if parallel processing should be performed. Such objects can be created by the SerialParam, MulticoreParam or SnowParam functions.	

xcmsSet

stopOnError	Logical specifying whether the feature detection call should stop on the first encountered error (the default), or whether feature detection is performed in all
	files regardless eventual failures for individual files in which case all errors are reported as warnings.
	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. By specifying a value of 0 the profile matrix generation can be skipped.

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, profBin

xcmsSet-class

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", ...).

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. The scan range provided in this slot represents the scans to which the whole raw data is subsetted.
- **.processHistory** Internal slot to be used to keep track of performed processing steps. This slot should not be directly accessed by the user.

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 = profStep(object), profparam=profinfo(object), mslevel = NULL, scanrange = NULL, rt=c("corrected", "r = bpparam()): 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. Argument BPPARAM allows to setup parallel processing.

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

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

progressCallback signature(object = "xcmsSet"): get the progressCallback slot

scanrange Getter method for the scanrange slot. See scanrange slot description above for more details.

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).

Author(s)

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

xcmsSource-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

xdata

LC-MS preprocessing result test data

Description

The 'xdata' variable represent the results from a 'xcms'-based pre-processing of an LC-MS untargeted metabolomics data set. The raw data files are provided in the 'faahKO' package. The pre-processing of this data set is described in detail in the *xcms* vignette of the 'xcms' package. [,xcmsRaw,logicalOrNumeric,missing,missing-method Subset an xcmsRaw object by scans

Description

Subset an xcmsRaw object by scans. The returned xcmsRaw object contains values for all scans specified with argument i. Note that the scanrange slot of the returned xcmsRaw will be c(1,length(object@scantime)) and hence not range(i).

Usage

S4 method for signature 'xcmsRaw,logicalOrNumeric,missing,missing' x[i, j, drop]

Arguments

х	The xcmsRaw object that should be sub-setted.
i	Integer or logical vector specifying the scans/spectra to which x should be sub- setted.
j	Not supported.
drop	Not supported.

Details

Only subsetting by scan index in increasing order or by a logical vector are supported. If not ordered, argument i is sorted automatically. Indices which are larger than the total number of scans are discarded.

Value

The sub-setted xcmsRaw object.

Author(s)

Johannes Rainer

See Also

split.xcmsRaw

Examples

```
## Load a test file
file <- system.file('cdf/K0/ko15.CDF', package = "faahKO")
xraw <- xcmsRaw(file, profstep = 0)
## The number of scans/spectra:
length(xraw@scantime)
## Subset the object to scans with a scan time from 3500 to 4000.
xsub <- xraw[xraw@scantime >= 3500 & xraw@scantime <= 4000]
range(xsub@scantime)
```

The number of scans: length(xsub@scantime) ## The number of values of the subset: length(xsub@env\$mz)

Index

```
* chromatographic peak refinement
        methods
    CleanPeaksParam, 38
    FilterIntensityParam, 94
    MergeNeighboringPeaksParam, 186
* classes
    xcmsEIC-class, 265
    xcmsFileSource-class, 266
    xcmsFragments-class, 268
    xcmsPeaks-class, 280
    xcmsRaw-class, 283
    xcmsSet-class, 288
    xcmsSource-class. 290
* core peak detection functions
    do_findChromPeaks_centWave, 48
    do_findChromPeaks_centWaveWithPredIsoROIs,
        52
    do_findChromPeaks_massifquant, 56
    do_findChromPeaks_matchedFilter,
        59
    do_findPeaks_MSW, 63
* core peak grouping algorithms
    do_groupChromPeaks_density, 64
    do_groupChromPeaks_nearest, 66
    do_groupPeaks_mzClust, 68
* core retention time correction algorithms
    do_adjustRtime_peakGroups, 45
* feature grouping methods
    groupFeatures-abundance-correlation,
        165
    groupFeatures-eic-similarity, 167
    groupFeatures-similar-rtime, 170
* file
    calibrate-methods, 31
    diffreport-methods, 43
    fillPeaks-methods, 84
    fillPeaks.chrom-methods, 85
    fillPeaks.MSW-methods, 86
    getEIC-methods, 146
```

getXcmsRaw-methods, 149 group.density, 151 group.mzClust, 152 group.nearest, 153 groupnames-methods, 172 peakTable-methods, 195 retcor.peakgroups-methods, 233 sampnames-methods, 235 verify.mzQuantM, 247 write.cdf-methods, 248 write.mzdata-methods, 249 write.mzQuantML-methods, 249 writeMzTab. 251 xcmsFileSource-class, 266 xcmsFragments, 267 xcmsRaw, 281 xcmsSet, 286 * functions to define bins breaks_on_binSize, 26 breaks_on_nBins, 28 * hplot image-methods, 176 levelplot-methods, 182 plot.xcmsEIC, 197 plotChrom-methods, 200 plotPeaks-methods, 211 plotRaw-methods, 213 plotrt-methods, 214 plotScan-methods, 215 plotSpec-methods, 215 plotSurf-methods, 216 plotTIC-methods, 217 * imputation functions imputeRowMin, 179 imputeRowMinRand, 180 * iplot plotChrom-methods, 200 plotSpec-methods, 215 plotSurf-methods, 216

```
plotTIC-methods, 217
                                                    group.mzClust, 152
* lockmass
                                                    group.nearest, 153
    AutoLockMass-methods, 20
                                                    groupnames-methods, 172
* manip
                                                    groupval-methods, 173
    AutoLockMass-methods, 20
                                                    loadRaw-methods, 183
    c-methods, 29
                                                    peakPlots-methods, 190
                                                    peakTable-methods, 195
    getPeaks-methods, 147
    getScan-methods, 148
                                                    plot.xcmsEIC, 197
    getSpec-methods, 148
                                                    plotChrom-methods, 200
                                                    plotEIC-methods, 208
    groupval-methods, 173
    medianFilter, 185
                                                    plotPeaks-methods, 211
    msn2xcmsRaw, 188
                                                    plotRaw-methods, 213
    profMedFilt-methods, 221
                                                    plotrt-methods, 214
    profMethod-methods, 221
                                                    plotScan-methods, 215
    profRange-methods, 222
                                                    plotSpec-methods, 215
    profStep-methods, 223
                                                    plotSurf-methods, 216
    retexp, 234
                                                    plotTIC-methods, 217
    specNoise, 240
                                                    profMedFilt-methods, 221
    specPeaks, 241
                                                    profMethod-methods, 221
    split.xcmsRaw, 242
                                                    profRange-methods, 222
    split.xcmsSet, 243
                                                    profStep-methods, 223
    stitch-methods, 244
                                                    rawEIC-methods, 226
* methods
                                                    rawMat-methods, 227
    absent-methods, 6
                                                    retcor-methods, 230
    AutoLockMass-methods, 20
                                                    retcor.obiwarp, 231
    calibrate-methods. 31
                                                    retcor.peakgroups-methods, 233
    collect-methods, 40
                                                    sampnames-methods, 235
    diffreport-methods, 43
                                                    specDist-methods, 237
    fillPeaks-methods, 84
                                                    specDist.cosine, 238
                                                    specDist.meanMZmatch, 239
    fillPeaks.chrom-methods, 85
    fillPeaks.MSW-methods, 86
                                                    specDist.peakCount-methods, 240
    findMZ, 122
                                                    stitch-methods, 244
    findneutral, 124
                                                    write.cdf-methods, 248
    findPeaks-methods, 125
                                                    write.mzdata-methods, 249
    findPeaks.addPredictedIsotopeFeatures-methodswrite.mzQuantML-methods, 249
        131
                                                    xcmsSource-methods, 291
    findPeaks.centWave-methods, 133
                                                * models
    findPeaks.centWaveWithPredictedIsotopeROIs-metbgds0
                                                * nonlinear
        135
    findPeaks.massifguant-methods, 138
                                                    SSgauss, 243
    findPeaks.MS1-methods, 142
                                                * peak detection functions for
                                                        chromatographic data
    getEIC-methods, 146
                                                    peaksWithCentWave, 191
    getPeaks-methods, 147
    getScan-methods, 148
                                                    peaksWithMatchedFilter, 194
                                                * peak detection methods
    getSpec-methods, 148
    getXcmsRaw-methods, 149
                                                    chromatographic-peak-detection, 35
    group-methods, 150
                                                    findChromPeaks-centWave, 99
    group.density, 151
                                                    findChromPeaks-centWaveWithPredIsoROIs,
```

105

findChromPeaks-massifquant, 109 findChromPeaks-matchedFilter, 116 findPeaks-MSW, 126 * peak grouping methods groupChromPeaks, 154 groupChromPeaks-density, 155 groupChromPeaks-mzClust, 159 groupChromPeaks-nearest, 162 * retention time correction methods adjustRtime. 7 adjustRtime-obiwarp, 7 adjustRtime-peakGroups, 13 [,XCMSnExp,ANY,ANY,ANY-method (filterFeatureDefinitions), 90 [,XChromatograms,ANY,ANY,ANY-method (XChromatograms), 252 [,xcmsRaw,logicalOrNumeric,missing,missing-method, 292 [,xcmsSet,ANY,ANY,ANY-method (xcmsSet-class), 288 [,xcmsSet-method(xcmsSet-class), 288 [[,XCMSnExp,ANY,ANY-method (filterFeatureDefinitions), 90 \$,xcmsSet-method(xcmsSet-class), 288 \$<-,xcmsSet-method (xcmsSet-class), 288</pre> absent (absent-methods), 6 absent, xcmsSet-method (absent-methods), 6 absent-methods, 6 absMz (groupChromPeaks-mzClust), 159 absMz,MzClustParam-method (groupChromPeaks-mzClust), 159 absMz.NearestPeaksParam-method (groupChromPeaks-nearest), 162 absMz<- (groupChromPeaks-mzClust), 159</pre> absMz<-,MzClustParam-method (groupChromPeaks-mzClust), 159 absMz<-,NearestPeaksParam-method (groupChromPeaks-nearest), 162 absRt (groupChromPeaks-nearest), 162 absRt,NearestPeaksParam-method (groupChromPeaks-nearest), 162 absRt<- (groupChromPeaks-nearest), 162 absRt<-,NearestPeaksParam-method (groupChromPeaks-nearest), 162 AbundanceSimilarityParam(), 74, 165, 166 addParams (findPeaks-MSW), 126

addParams, MSWParam-method (findPeaks-MSW), 126 addParams<- (findPeaks-MSW), 126 addParams<-,MSWParam-method (findPeaks-MSW), 126 adjustedRtime, 11, 16 adjustedRtime (XCMSnExp-class), 269 adjustedRtime,MsFeatureData-method (XCMSnExp-class), 269 adjustedRtime,XCMSnExp-method (XCMSnExp-class), 269 adjustedRtime<- (XCMSnExp-class), 269 adjustedRtime<-,MsFeatureData-method (XCMSnExp-class), 269 adjustedRtime<-,XCMSnExp-method (XCMSnExp-class), 269 adjustRtime, 7, 12, 17, 199, 269, 270, 278, 279 adjustRtime(), 19 adjustRtime,OnDiskMSnExp,ObiwarpParam-method (adjustRtime-obiwarp), 7 adjustRtime,XCMSnExp,ObiwarpParam-method (adjustRtime-obiwarp), 7 adjustRtime,XCMSnExp,PeakGroupsParam-method (adjustRtime-peakGroups), 13 adjustRtime-obiwarp, 7 adjustRtime-peakGroups, 13 adjustRtimePeakGroups, 16 adjustRtimePeakGroups (adjustRtime-peakGroups), 13 alignRt(), 167, 168 ampTh (findPeaks-MSW), 126 ampTh,MSWParam-method(findPeaks-MSW), 126 ampTh<- (findPeaks-MSW), 126 ampTh<-,MSWParam-method</pre> (findPeaks-MSW), 126 applyAdjustedRtime, 18, 278 applyAdjustedRtime(), 72, 93 array, 281 as, 279 AutoLockMass (AutoLockMass-methods), 20 AutoLockMass,xcmsRaw-method (AutoLockMass-methods), 20 AutoLockMass-methods, 20 baseValue

(findChromPeaks-matchedFilter), 116

baseValue, MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 baseValue<-(findChromPeaks-matchedFilter), 116 baseValue<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 bin, 21 bin,XCMSnExp-method,21 binSize (findChromPeaks-matchedFilter), 116 binSize,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 binSize,ObiwarpParam-method (adjustRtime-obiwarp), 7 binSize, PeakDensityParam-method (groupChromPeaks-density), 155 binSize<-(findChromPeaks-matchedFilter), 116 binSize<-.MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 binSize<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 binSize <-, PeakDensityParam-method (groupChromPeaks-density), 155 binYonX, 23, 27, 28, 61, 62, 120, 220, 264 bpparam, 85, 103, 108, 114, 119, 129, 276 bpparam(), 70, 95, 96, 185, 187, 228 breaks_on_binSize, 24, 26, 28 breaks_on_nBins, 24, 27, 28 bw (groupChromPeaks-density), 155 bw, PeakDensityParam-method (groupChromPeaks-density), 155 bw<- (groupChromPeaks-density), 155</pre> bw<-,PeakDensityParam-method</pre> (groupChromPeaks-density), 155

c, 289

c, c-methods(c-methods), 29 c-methods, 29 c.XCMSnExp(XCMSnExp-class), 269 c.xcmsSet(c-methods), 29 CalibrantMassParam (CalibrantMassParam-class), 29 CalibrantMassParam-class, 29 calibrate (calibrate-methods), 31 calibrate,XCMSnExp-method (CalibrantMassParam-class), 29 calibrate,xcmsSet-method (calibrate-methods), 31 calibrate-methods, 31 centerSample (adjustRtime-obiwarp), 7 centerSample,ObiwarpParam-method (adjustRtime-obiwarp), 7 centerSample<- (adjustRtime-obiwarp), 7 centerSample<-,ObiwarpParam-method (adjustRtime-obiwarp), 7 centWave, 35, 51, 55, 96, 97, 108, 135, 140, 192, 193 centWave (findChromPeaks-centWave), 99 CentWaveParam, 96, 105, 109, 122, 145, 219, 276 CentWaveParam (findChromPeaks-centWave), 99 CentWaveParam-class (findChromPeaks-centWave), 99 CentWavePredIsoParam, 276 CentWavePredIsoParam (findChromPeaks-centWaveWithPredIsoROIs), 105 CentWavePredIsoParam-class (findChromPeaks-centWaveWithPredIsoROIs), 105 centWaveWithPredIsoROIs, 35 centWaveWithPredIsoROIs (findChromPeaks-centWaveWithPredIsoROIs), 105 character, 266 CharacterList(), 229 checkBack(findChromPeaks-massifquant), 109 checkBack,MassifquantParam-method (findChromPeaks-massifquant), 109 checkBack<-(findChromPeaks-massifquant), 109 checkBack<-,MassifguantParam-method</pre> (findChromPeaks-massifquant), 109 Chromatogram, 34, 96, 98, 174, 258, 259, 278, 279

chromatogram, 279 chromatogram (chromatogram, XCMSnExp-method), 32 Chromatogram(), 42, 87, 230, 252 chromatogram(), 76, 259, 261 chromatogram, XCMSnExp-method, 32 chromatographic-peak-detection, 35 ChromPeakAreaParam (FillChromPeaksParam-class), 80 ChromPeakAreaParam-class (FillChromPeaksParam-class), 80 chromPeakData (XCMSnExp-class), 269 chromPeakData(), 80, 121, 122 chromPeakData,MsFeatureData-method (XCMSnExp-class), 269 chromPeakData,XChromatogram-method (XChromatograms), 252 chromPeakData,XChromatograms-method (XChromatograms), 252 chromPeakData, XCMSnExp-method (XCMSnExp-class), 269 chromPeakData<- (XCMSnExp-class), 269 chromPeakData<-,MsFeatureData-method (XCMSnExp-class), 269 chromPeakData<-,XChromatogram-method</pre> (XChromatograms), 252 chromPeakData<-,XCMSnExp-method (XCMSnExp-class), 269 chromPeaks, 11, 16, 175, 205, 224 chromPeaks (XCMSnExp-class), 269 chromPeaks(), 37, 80, 83, 94, 95, 121 chromPeaks,MsFeatureData-method (XCMSnExp-class), 269 chromPeaks, XChromatogram-method (XChromatograms), 252 chromPeaks, XChromatograms-method (XChromatograms), 252 chromPeaks,XCMSnExp-method (XCMSnExp-class), 269 chromPeaks<- (XCMSnExp-class), 269 chromPeaks<-,MsFeatureData-method (XCMSnExp-class), 269 chromPeaks<-,XChromatogram-method (XChromatograms), 252 chromPeaks<-,XCMSnExp-method (XCMSnExp-class), 269 chromPeakSpectra, 36, 279

chromPeakSpectra(), 77, 78 class:Param (GenericParam-class), 145 clean, 21 clean(), 229 clean, XCMSnExp-method (bin,XCMSnExp-method), 21 CleanPeaksParam, 38, 95, 188 CleanPeaksParam-class (CleanPeaksParam), 38 coerce, MChromatograms, XChromatograms-method (XChromatograms), 252 collect, 267, 268 collect (collect-methods), 40 collect, xcmsFragments-method (collect-methods), 40 collect, xcmsRaw-method (collect-methods), 40 collect-methods, 40 compareChromatograms(), 41, 167, 168 consecMissedLimit (findChromPeaks-massifquant), 109 consecMissedLimit,MassifquantParam-method (findChromPeaks-massifquant), 109 consecMissedLimit<-(findChromPeaks-massifquant), 109 consecMissedLimit<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 cor(), 42, 168 correlate (correlate, Chromatogram, Chromatogram-method), 41 correlate, Chromatogram, Chromatogram-method, 41 correlate, MChromatograms, MChromatograms-method (correlate, Chromatogram, Chromatogram-method), 41 correlate, MChromatograms, missing-method (correlate, Chromatogram, Chromatogram-method), 41 criticalValue (findChromPeaks-massifquant), 109 criticalValue,MassifquantParam-method (findChromPeaks-massifquant),

109 criticalValue<-(findChromPeaks-massifquant), 109 criticalValue<-,MassifquantParam-method (findChromPeaks-massifquant), 109

DataFrame(), 259 deepCopy (xcmsRaw), 281 deepCopy, xcmsRaw-method (xcmsRaw), 281 density, 151 diffreport, 6, 196, 289 diffreport (diffreport-methods), 43 diffreport, xcmsSet-method (diffreport-methods), 43 diffreport-methods, 43 dirname, 45 dirname, OnDiskMSnExp-method (dirname), 45 dirname<-,OnDiskMSnExp-method (dirname), 45 distance (findChromPeaks-matchedFilter), 116 distance, MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 distance<-(findChromPeaks-matchedFilter), 116 distance<-,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 distFun (adjustRtime-obiwarp), 7 distFun,ObiwarpParam-method (adjustRtime-obiwarp), 7 distFun<- (adjustRtime-obiwarp), 7</pre> distFun<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 do_adjustRtime_peakGroups, 17, 45 do_findChromPeaks_addPredIsoROIs (do_findChromPeaks_centWaveWithPredIsoROIs), (groupFeatures-eic-similarity), 52 do_findChromPeaks_centWave, 48, 56, 58, 59, 62, 64, 104, 113

do_findChromPeaks_centWaveWithPredIsoROIs, 51, 52, 59, 62, 64, 109, 138

do_findChromPeaks_massifquant, 51, 56, 56, 62, 64, 115 do_findChromPeaks_matchedFilter, 51, 56, 59, 59, 64, 120, 140, 142, 247 do_findPeaks_MSW, 51, 56, 59, 62, 63, 130, 145 do_groupChromPeaks_density, 64, 67, 69, 151 do_groupChromPeaks_density(), 158 do_groupChromPeaks_nearest, 66, 66, 69 do_groupChromPeaks_nearest(), 164 do_groupPeaks_mzClust, 66, 67, 68 do_groupPeaks_mzClust(), 161 dropAdjustedRtime (XCMSnExp-class), 269 dropAdjustedRtime(), 19 dropAdjustedRtime,MsFeatureData-method (XCMSnExp-class), 269 dropAdjustedRtime,XCMSnExp-method (XCMSnExp-class), 269 dropChromPeaks (XCMSnExp-class), 269 dropChromPeaks,MsFeatureData-method (XCMSnExp-class), 269 dropChromPeaks,XCMSnExp-method (XCMSnExp-class), 269 dropFeatureDefinitions (XCMSnExp-class), 269 dropFeatureDefinitions,MsFeatureData-method (XCMSnExp-class), 269 dropFeatureDefinitions,XChromatograms-method (XChromatograms), 252 dropFeatureDefinitions,XCMSnExp-method (XCMSnExp-class), 269 dropFilledChromPeaks (XCMSnExp-class), 269 dropFilledChromPeaks(), 259 dropFilledChromPeaks,XChromatogram-method (XChromatograms), 252 dropFilledChromPeaks,XChromatograms-method (XChromatograms), 252 dropFilledChromPeaks,XCMSnExp-method (XCMSnExp-class), 269 EicSimilarityParam

167

167

EicSimilarityParam(), 74

EicSimilarityParam-class

(groupFeatures-eic-similarity),

estimatePrecursorIntensity, 69 etg. 70 expandMz (FillChromPeaksParam-class), 80 expandMz,FillChromPeaksParam-method (FillChromPeaksParam-class), 80 expandMz<- (FillChromPeaksParam-class),</pre> 80 expandMz<-,FillChromPeaksParam-method</pre> (FillChromPeaksParam-class), 80 expandRt (FillChromPeaksParam-class), 80 expandRt,FillChromPeaksParam-method (FillChromPeaksParam-class), 80 expandRt<- (FillChromPeaksParam-class),</pre> 80 expandRt<-,FillChromPeaksParam-method</pre> (FillChromPeaksParam-class), 80 exportMetaboAnalyst,71 extractMsData (extractMsData, OnDiskMSnExp-method), 72 extractMsData(), 211 extractMsData,OnDiskMSnExp-method,72 extractMsData,XCMSnExp-method (extractMsData,OnDiskMSnExp-method), 72 extraPeaks (adjustRtime-peakGroups), 13 extraPeaks, PeakGroupsParam-method (adjustRtime-peakGroups), 13 extraPeaks<- (adjustRtime-peakGroups),</pre> 13 extraPeaks<-, PeakGroupsParam-method (adjustRtime-peakGroups), 13 faahko_sub (XCMSnExp-class), 269 factorDiag(adjustRtime-obiwarp), 7 factorDiag,ObiwarpParam-method (adjustRtime-obiwarp), 7 factorDiag<- (adjustRtime-obiwarp), 7</pre> factorDiag<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 factorGap (adjustRtime-obiwarp), 7 factorGap,ObiwarpParam-method (adjustRtime-obiwarp), 7 factorGap<- (adjustRtime-obiwarp), 7</pre> factorGap<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 family (adjustRtime-peakGroups), 13 family,PeakGroupsParam-method (adjustRtime-peakGroups), 13

family<- (adjustRtime-peakGroups), 13</pre> family<-,PeakGroupsParam-method</pre> (adjustRtime-peakGroups), 13 feature-grouping, 74 featureArea, 83 featureArea (XCMSnExp-class), 269 featureChromatograms, 75, 155, 225, 279 featureDefinitions, 154, 155, 224, 225, 279 featureDefinitions (XCMSnExp-class), 269 featureDefinitions(), 78, 80, 157, 158, 161, 164, 170, 190 featureDefinitions,MsFeatureData-method (XCMSnExp-class), 269 featureDefinitions,XChromatograms-method (XChromatograms), 252 featureDefinitions,XCMSnExp-method (XCMSnExp-class), 269 featureDefinitions<- (XCMSnExp-class),</pre> 269 featureDefinitions<-,MsFeatureData-method (XCMSnExp-class), 269 featureDefinitions<-,XCMSnExp-method</pre> (XCMSnExp-class), 269 featureGroups(), 209 featureGroups, XCMSnExp-method (feature-grouping), 74 featureGroups<-,XCMSnExp-method</pre> (feature-grouping), 74 featureSpectra, 77, 279 featureSummary, 78, 270, 278, 279 featureValues, 224, 278 featureValues (quantify, XCMSnExp-method), 224 featureValues(), 71, 72, 83, 158, 161, 164, 166 featureValues, XChromatograms-method (XChromatograms), 252 featureValues, XCMSnExp-method (quantify,XCMSnExp-method), 224 fileIndex (ProcessHistory-class), 217 fileIndex, ProcessHistory-method (ProcessHistory-class), 217 filepaths (xcmsSet-class), 288 filepaths, xcmsSet-method (xcmsSet-class), 288 filepaths<- (xcmsSet-class), 288 filepaths<-,xcmsSet-method</pre>

(xcmsSet-class), 288 filterColumnsKeepTop (filterColumnsIntensityAbove,MChromatograms-method fillChromPeaks, 225, 271, 277, 279 87 fillChromPeaks filterColumnsKeepTop(), 76 (FillChromPeaksParam-class), 80 $fill Chrom {\tt Peaks, XCMSnExp, Chrom {\tt PeakAreaParam-mefbbd} erColumns {\tt KeepTop, MChromatograms-method}}$ (FillChromPeaksParam-class), 80 (filterColumnsIntensityAbove,MChromatograms-method fillChromPeaks,XCMSnExp,FillChromPeaksParam-method 87 (FillChromPeaksParam-class), 80 filterColumnsKeepTop,XChromatograms-method (filterColumnsIntensityAbove,MChromatograms-method fillChromPeaks,XCMSnExp,missing-method 87 (FillChromPeaksParam-class), 80 filterFeatureDefinitions, 90 FillChromPeaksParam (FillChromPeaksParam-class), 80 filterFile,XCMSnExp-method FillChromPeaksParam-class. 80 (filterFeatureDefinitions), 90 fillPeaks, 6, 43, 86, 87, 288, 289 filterIntensity(), 229 FilterIntensityParam, 39, 94, 188 fillPeaks (fillPeaks-methods), 84 fillPeaks,xcmsSet-method FilterIntensityParam-class (FilterIntensityParam), 94 (fillPeaks-methods), 84 fillPeaks-methods, 84 filterMsLevel,XCMSnExp-method fillPeaks.chrom, 87 (filterFeatureDefinitions), 90 fillPeaks.chrom filterMz,XChromatogram-method (XChromatograms), 252 (fillPeaks.chrom-methods), 85 fillPeaks.chrom,xcmsSet-method filterMz,XChromatograms-method (XChromatograms), 252 (fillPeaks.chrom-methods), 85 fillPeaks.chrom-methods, 85 filterMz,XCMSnExp-method fillPeaks.MSW (fillPeaks.MSW-methods), (filterFeatureDefinitions), 90 86 filterRt,XChromatogram-method fillPeaks.MSW,xcmsSet-method (XChromatograms), 252 (fillPeaks.MSW-methods), 86 filterRt,XChromatograms-method fillPeaks.MSW-methods, 86 (XChromatograms), 252 filterRt,XCMSnExp-method filterAcquisitionNum, 21 (filterFeatureDefinitions), 90 filterAcquisitionNum,XCMSnExp-method (bin, XCMSnExp-method), 21 findChromPeaks, 16, 217, 270, 277, 279 findChromPeaks filterChromPeaks (XChromatograms), 252 filterChromPeaks,XChromatogram-method (chromatographic-peak-detection), (XChromatograms), 252 35 filterChromPeaks,XChromatograms-method findChromPeaks(), 121, 122, 157, 160, 164, (XChromatograms), 252 184 filterChromPeaks,XCMSnExp-method findChromPeaks,Chromatogram,CentWaveParam-method, (filterFeatureDefinitions), 90 96 filterColumnsIntensityAbove findChromPeaks, Chromatogram, MatchedFilterParam-method, (filterColumnsIntensityAbove,MChromatograms-m@8hod), findChromPeaks, MChromatograms, CentWaveParam-method 87 (findChromPeaks, Chromatogram, CentWaveParam-method), filterColumnsIntensityAbove(), 259 filterColumnsIntensityAbove,MChromatograms-method, 96 findChromPeaks, MChromatograms, MatchedFilterParam-method 87 filterColumnsIntensityAbove,XChromatograms-method (findChromPeaks, Chromatogram, CentWaveParam-method), (filterColumnsIntensityAbove,MChromatograms-m@6hod), 87 findChromPeaks, OnDiskMSnExp, CentWaveParam-method

(findChromPeaks-centWave), 99 findPeaks.addPredictedIsotopeFeatures,xcmsRaw-method findChromPeaks,OnDiskMSnExp,CentWavePredIsoParam-meth@GindPeaks.addPredictedIsotopeFeatures-methods), (findChromPeaks-centWaveWithPredIsoROIs), 131 105 findPeaks.addPredictedIsotopeFeatures-methods, findChromPeaks,OnDiskMSnExp,MassifquantParam-method 131 (findChromPeaks-massifquant), findPeaks.centWave, 43, 104, 109, 126, 133, 109 138 findChromPeaks,OnDiskMSnExp,MatchedFilterParafiimdReakds.centWave (findChromPeaks-matchedFilter), (findPeaks.centWave-methods), 116 133 findChromPeaks,OnDiskMSnExp,MSWParam-method findPeaks.centWave,xcmsRaw-method (findPeaks-MSW), 126 (findPeaks.centWave-methods), findChromPeaks, XCMSnExp, Param-method 133 (XCMSnExp-class), 269 findPeaks.centWave-methods, 133 findChromPeaks-centWave, 99, 261 findPeaks.centWaveWithPredictedIsotopeROIs, findChromPeaks-centWaveWithPredIsoROIs, 126 105 findPeaks.centWaveWithPredictedIsotopeROIs findChromPeaks-Chromatogram-CentWaveParam, (findPeaks.centWaveWithPredictedIsotopeROIs-method 260 135 findChromPeaks-Chromatogram-CentWaveParam findPeaks.centWaveWithPredictedIsotopeROIs,xcmsRaw-method (findChromPeaks, Chromatogram, CentWaveParam-metfixeddPeaks.centWaveWithPredictedIsotopeROIs-method 135 96 findChromPeaks-massifquant, 109 findPeaks.centWaveWithPredictedIsotopeROIs-methods, findChromPeaks-matchedFilter, 116 135 findChromPeaksIsolationWindow, 121 findPeaks.massifquant, 115 findChromPeaksIsolationWindow(), 229 findPeaks.massifquant (findPeaks.massifquant-methods), findMZ, 122, 125 findMZ,xcmsFragments-method (findMZ), 138 122 findPeaks.massifquant,xcmsRaw-method (findPeaks.massifquant-methods), findmzROI (xcmsRaw-class), 283 138 findmzROI, xcmsRaw-method findPeaks.massifquant-methods, 138 (xcmsRaw-class), 283 findPeaks.matchedFilter, 120, 126, 287 findneutral, *123*, 124 findPeaks.matchedFilter findneutral,xcmsFragments-method (findPeaks.matchedFilter,xcmsRaw-method), (findneutral), 124 140 findPeaks, 35, 104, 109, 115, 120, 130, 147, 191, 211, 212, 221, 236, 280, 281, findPeaks.matchedFilter,xcmsRaw-method, 284, 287 140 findPeaks (findPeaks-methods), 125 findPeaks.MS1 (findPeaks.MS1-methods), 142 findPeaks,xcmsRaw-method (findPeaks-methods), 125 findPeaks.MS1,xcmsRaw-method (findPeaks.MS1-methods), 142 findPeaks-methods, 125 findPeaks.MS1-methods, 142 findPeaks-MSW, 126 findPeaks.MSW, 130 findPeaks.addPredictedIsotopeFeatures, findPeaks.MSW 126.138 findPeaks.addPredictedIsotopeFeatures (findPeaks.MSW, xcmsRaw-method), (findPeaks.addPredictedIsotopeFeatures-methods)44 131 findPeaks.MSW, xcmsRaw-method, 144

firstBaselineCheck (findChromPeaks-centWave), 99 firstBaselineCheck,CentWaveParam-method (findChromPeaks-centWave), 99 firstBaselineCheck<-(findChromPeaks-centWave), 99 firstBaselineCheck<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 fitgauss (findChromPeaks-centWave), 99 fitgauss, CentWaveParam-method (findChromPeaks-centWave), 99 fitgauss,MassifquantParam-method (findChromPeaks-massifquant), 109 fitgauss<- (findChromPeaks-centWave), 99 fitgauss<-,CentWaveParam-method (findChromPeaks-centWave), 99 fitgauss<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 fixedMz (FillChromPeaksParam-class), 80 fixedRt (FillChromPeaksParam-class), 80 format(), 71fwhm (findChromPeaks-matchedFilter), 116 fwhm,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 fwhm<- (findChromPeaks-matchedFilter),</pre> 116 fwhm<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 gapExtend (adjustRtime-obiwarp), 7 gapExtend,ObiwarpParam-method (adjustRtime-obiwarp), 7

getEIC, xcmsRaw-method (getEIC-methods), 146 getEIC, xcmsSet-method (getEIC-methods), 146 getEIC-methods, 146 getMsnScan (getScan-methods), 148 getMsnScan, xcmsRaw-method (getScan-methods), 148 getPeaks, 85-87, 247, 284 getPeaks (getPeaks-methods), 147 getPeaks, xcmsRaw-method (getPeaks-methods), 147 getPeaks-methods, 147 getScan, 149, 284 getScan (getScan-methods), 148 getScan, xcmsRaw-method (getScan-methods), 148 getScan-methods, 148 getSpec, 148, 241, 242, 284 getSpec (getSpec-methods), 148 getSpec, xcmsRaw-method (getSpec-methods), 148 getSpec-methods, 148 getXcmsRaw, 289 getXcmsRaw (getXcmsRaw-methods), 149 getXcmsRaw, xcmsSet-method (getXcmsRaw-methods), 149 getXcmsRaw-methods, 149 group, 6, 16, 155, 288, 289 group (group-methods), 150 group(), 161 group,xcmsSet-method (group-methods), 150 group-methods, 150 group.density, 150, 151, 153 group.density(), 158 group.density,xcmsSet-method (group.density), 151 group.mzClust, 150, 152, 153 group.mzClust(), 161 group.mzClust,xcmsSet-method (group.mzClust), 152 group.nearest, 150, 153 group.nearest,xcmsSet-method (group.nearest), 153 groupChromPeaks, 16, 17, 154, 158, 161, 164, 261, 278, 279 groupChromPeaks(), 74, 83

```
groupChromPeaks, XChromatograms, PeakDensityPargmoupvalder, 173
                                               groupval, xcmsSet-method
        (XChromatograms), 252
groupChromPeaks,XCMSnExp,MzClustParam-method
                                                       (groupval-methods), 173
        (groupChromPeaks-mzClust), 159
                                               groupval-methods, 173
groupChromPeaks,XCMSnExp,NearestPeaksParam-method
        (groupChromPeaks-nearest), 162
                                               hasAdjustedRtime (XCMSnExp-class), 269
groupChromPeaks,XCMSnExp,PeakDensityParam-methedAdjustedRtime,MsFeatureData-method
        (groupChromPeaks-density), 155
                                                       (XCMSnExp-class), 269
groupChromPeaks-density, 155
                                               hasAdjustedRtime,OnDiskMSnExp-method
groupChromPeaks-mzClust, 159
                                                       (XCMSnExp-class), 269
groupChromPeaks-nearest, 162
                                               hasAdjustedRtime,XCMSnExp-method
groupFeatures(), 210
                                                       (XCMSnExp-class), 269
groupFeatures, XCMSnExp, AbundanceSimilarityParkans@ketbonPeaks (XCMSnExp-class), 269
        (groupFeatures-abundance-correlation), hasChromPeaks, MsFeatureData-method
        165
                                                       (XCMSnExp-class), 269
groupFeatures,XCMSnExp,EicSimilarityParam-method
        (groupFeatures-eic-similarity),
                                                       (XChromatograms), 252
        167
                                               hasChromPeaks,XChromatograms-method
groupFeatures,XCMSnExp,SimilarRtimeParam-method
                                                       (XChromatograms), 252
        (groupFeatures-similar-rtime),
                                               hasChromPeaks, XCMSnExp-method
        170
                                                       (XCMSnExp-class), 269
groupFeatures-abundance-correlation,
                                               hasFeatures, 225
                                               hasFeatures(XCMSnExp-class), 269
        165
groupFeatures-eic-similarity, 167
                                               hasFeatures,MsFeatureData-method
groupFeatures-similar-rtime, 170
                                                       (XCMSnExp-class), 269
groupidx (xcmsSet-class), 288
                                               hasFeatures,XChromatograms-method
groupidx, xcmsSet-method
                                                       (XChromatograms), 252
        (xcmsSet-class), 288
                                               hasFeatures,XCMSnExp-method
groupidx<- (xcmsSet-class), 288</pre>
                                                       (XCMSnExp-class), 269
groupidx<-,xcmsSet-method</pre>
                                               hasFilledChromPeaks(XCMSnExp-class),
        (xcmsSet-class), 288
                                                       269
groupnames, 72, 265, 289
                                               hasFilledChromPeaks,XChromatograms-method
groupnames (groupnames-methods), 172
                                                       (XChromatograms), 252
                                               hasFilledChromPeaks,XCMSnExp-method
groupnames, xcmsEIC-method
                                                       (XCMSnExp-class), 269
        (groupnames-methods), 172
groupnames, XCMSnExp-method, 171
                                               highlightChromPeaks, 35, 174, 208, 278
groupnames, xcmsSet-method
        (groupnames-methods), 172
                                               identifyMajorPeaks, 127, 129, 144
groupnames-methods, 172
                                               image, 285
groupOverlaps, 173
                                               image,xcmsRaw-method (image-methods),
groups (xcmsSet-class), 288
                                                       176
groups,xcmsSet-method(xcmsSet-class),
                                               image-methods, 176
        288
                                               impute,MatchedFilterParam-method
groups<- (xcmsSet-class), 288</pre>
                                                       (findChromPeaks-matchedFilter),
groups<-,xcmsSet-method</pre>
                                                       116
                                               impute<-</pre>
        (xcmsSet-class), 288
groupSimilarityMatrix(), 167, 168
                                                       (findChromPeaks-matchedFilter),
groupval, 195, 225, 289
                                                       116
```

impute<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 imputeLinInterpol, 25, 60-62, 119, 120, 177, 219, 220, 264, 275 imputeRowMin, 179, 181 imputeRowMinRand, 179, 180 index(findChromPeaks-matchedFilter), 116 index,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 index<- (findChromPeaks-matchedFilter),</pre> 116 index<-.MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 initPenalty (adjustRtime-obiwarp), 7 initPenalty,ObiwarpParam-method (adjustRtime-obiwarp), 7 initPenalty<- (adjustRtime-obiwarp), 7</pre> initPenalty<-,ObiwarpParam-method (adjustRtime-obiwarp), 7 integrate, CentWaveParam-method (findChromPeaks-centWave), 99 integrate, MassifquantParam-method (findChromPeaks-massifquant), 109 integrate<- (findChromPeaks-centWave),</pre> 99 integrate<-,CentWaveParam-method (findChromPeaks-centWave), 99 integrate<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 intensity, XCMSnExp-method (XCMSnExp-class), 269 isCalibrated (CalibrantMassParam-class), 29 isolationWindowTargetMz (isolationWindowTargetMz,OnDiskMSnExp^M和E的点面以antParam-class 182 isolationWindowTargetMz(), 121 isolationWindowTargetMz,OnDiskMSnExp-method, matchedFilter, 35, 62, 98, 142, 194, 195 182 kNN (groupChromPeaks-nearest), 162

kNN,NearestPeaksParam-method (groupChromPeaks-nearest), 162

kNN<- (groupChromPeaks-nearest), 162 kNN<-,NearestPeaksParam-method (groupChromPeaks-nearest), 162 lattice::level.colors, 211 levelplot, 285 levelplot (xcmsRaw-class), 283 levelplot,xcmsRaw-method (levelplot-methods), 182 levelplot.xcmsSet-method (levelplot-methods), 182 levelplot-methods, 182 loadRaw(loadRaw-methods), 183 loadRaw,xcmsFileSource-method (loadRaw-methods), 183 loadRaw,xcmsSource-method (loadRaw-methods), 183 loadRaw-methods, 183 localAlignment (adjustRtime-obiwarp), 7 localAlignment,ObiwarpParam-method (adjustRtime-obiwarp), 7 localAlignment<- (adjustRtime-obiwarp),</pre> localAlignment<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 loess, 15, 46, 233

makeacqNum (stitch-methods), 244 makeacqNum, xcmsRaw-method (stitch-methods), 244 manualChromPeaks, 35, 184 manualFeatures (manualChromPeaks), 184 massifquant, 35, 59 massifquant (findChromPeaks-massifquant), 109 MassifquantParam, 276 MassifquantParam (findChromPeaks-massifquant), 109 (findChromPeaks-massifquant), 109 matchedFilter (findChromPeaks-matchedFilter), 116 matchedFilter(), 83

MatchedFilterParam, 98, 276

MatchedFilterParam MChromatograms, 33, 96, 98, 174 (findChromPeaks-matchedFilter), MChromatograms(), 41, 42, 87, 88, 201, 202, 116 230, 252, 258, 259, 261 MatchedFilterParam-class medianFilter, 185, 221 (findChromPeaks-matchedFilter), MergeNeighboringPeaksParam, 39, 95, 186 116 MergeNeighboringPeaksParam(), 184, 260 matplot, 175 MergeNeighboringPeaksParam-class matrix, 281 (MergeNeighboringPeaksParam), max, MatchedFilterParam-method 186 (findChromPeaks-matchedFilter), minFraction (groupChromPeaks-density), 155 116 max<- (findChromPeaks-matchedFilter),</pre> minFraction,MzClustParam-method 116 (groupChromPeaks-mzClust), 159 max<-,MatchedFilterParam-method</pre> minFraction,PeakDensityParam-method (groupChromPeaks-density), 155 (findChromPeaks-matchedFilter), 116 minFraction.PeakGroupsParam-method (adjustRtime-peakGroups), 13 maxCharge (findChromPeaks-centWaveWithPredIsoROInshFraction<-105 (groupChromPeaks-density), 155 maxCharge.CentWavePredIsoParam-method minFraction<-,MzClustParam-method</pre> (findChromPeaks-centWaveWithPredIsoROIs), (groupChromPeaks-mzClust), 159 105 minFraction<-,PeakDensityParam-method</pre> (groupChromPeaks-density), 155 maxCharge<-(findChromPeaks-centWaveWithPredIsoROIns), Fraction <-, PeakGroupsParam-method 105 (adjustRtime-peakGroups), 13 minNoiseLevel (findPeaks-MSW), 126 maxCharge<-,CentWavePredIsoParam-method</pre> (findChromPeaks-centWaveWithPredIsoROInshNoiseLevel,MSWParam-method (findPeaks-MSW), 126 105 maxFeatures(groupChromPeaks-density), minNoiseLevel<- (findPeaks-MSW), 126</pre> 155 minNoiseLevel<-,MSWParam-method</pre> maxFeatures,PeakDensityParam-method (findPeaks-MSW), 126 (groupChromPeaks-density), 155 minSamples (groupChromPeaks-density), maxFeatures<-155 (groupChromPeaks-density), 155 minSamples, MzClustParam-method (groupChromPeaks-mzClust), 159 maxFeatures<-,PeakDensityParam-method</pre> (groupChromPeaks-density), 155 minSamples, PeakDensityParam-method maxIso (groupChromPeaks-density), 155 (findChromPeaks-centWaveWithPredIsoROInsin,Samples<- (groupChromPeaks-density), 155 105 minSamples<-,MzClustParam-method</pre> maxIso,CentWavePredIsoParam-method (findChromPeaks-centWaveWithPredIsoROIs), (groupChromPeaks-mzClust), 159 105 minSamples<-,PeakDensityParam-method</pre> (groupChromPeaks-density), 155 maxIso<-(findChromPeaks-centWaveWithPredIsoRO**Ids**, eatures::groupFeatures(), 74 105 MsFeatures::SimilarRtimeParam(), 170 maxIso<-,CentWavePredIsoParam-method</pre> mslevel (xcmsSet-class), 288 (findChromPeaks-centWaveWithPredIsoROIns),evel,xcmsRaw-method (xcmsRaw-class), 105 283

mslevel,xcmsSet-method(xcmsSet-class), 288 msLevel, XProcessHistory-method (ProcessHistory-class), 217 msn2xcmsRaw, 188 MSnbase::Chromatogram(), 41, 42 MSnExp, 271, 279 MSpectra, 37, 78 MSpectra(), 229 MSW, 35, 64, 145 MSW (findPeaks-MSW), 126 MSWParam, 276 MSWParam (findPeaks-MSW), 126 MSWParam-class (findPeaks-MSW), 126 MulticoreParam, 286 mz,CalibrantMassParam (CalibrantMassParam-class), 29 mz, XCMSnExp-method (XCMSnExp-class), 269 mzCenterFun (findChromPeaks-centWave), 99 mzCenterFun,CentWaveParam-method (findChromPeaks-centWave), 99 mzCenterFun,MassifquantParam-method (findChromPeaks-massifquant), 109 mzCenterFun<-(findChromPeaks-centWave), 99 mzCenterFun<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 mzCenterFun<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 MzClustParam, 161 MzClustParam (groupChromPeaks-mzClust), 159 MzClustParam-class (groupChromPeaks-mzClust), 159 mzdiff(findChromPeaks-centWave), 99 mzdiff,CentWaveParam-method (findChromPeaks-centWave), 99 mzdiff,MassifquantParam-method (findChromPeaks-massifquant), 109 mzdiff,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 mzdiff<- (findChromPeaks-centWave), 99</pre>

mzdiff<-,CentWaveParam-method</pre>

(findChromPeaks-centWave), 99 mzdiff<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 mzdiff<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 mzIntervalExtension (findChromPeaks-centWaveWithPredIsoROIs), 105 mzIntervalExtension,CentWavePredIsoParam-method (findChromPeaks-centWaveWithPredIsoROIs), 105 mzIntervalExtension<-</pre> (findChromPeaks-centWaveWithPredIsoROIs), 105 mzIntervalExtension<-,CentWavePredIsoParam-method</pre> (findChromPeaks-centWaveWithPredIsoROIs), 105 mzR::writeMSData(), 251 mzrange (xcmsEIC-class), 265 mzrange,xcmsEIC-method(xcmsEIC-class), 265 mzVsRtBalance (groupChromPeaks-nearest), 162 mzVsRtBalance,NearestPeaksParam-method (groupChromPeaks-nearest), 162 mzVsRtBalance<-(groupChromPeaks-nearest), 162 mzVsRtBalance<-,NearestPeaksParam-method (groupChromPeaks-nearest), 162 nearbyPeak (findPeaks-MSW), 126 nearbyPeak,MSWParam-method (findPeaks-MSW), 126 nearbyPeak<- (findPeaks-MSW), 126</pre> nearbyPeak<-,MSWParam-method</pre> (findPeaks-MSW), 126 NearestPeaksParam (groupChromPeaks-nearest), 162 NearestPeaksParam-class (groupChromPeaks-nearest), 162 nls, 244 noise(findChromPeaks-centWave), 99 noise, CentWaveParam-method (findChromPeaks-centWave), 99 noise, MassifquantParam-method

(findChromPeaks-massifquant), 109

noise<- (findChromPeaks-centWave), 99
noise<-,CentWaveParam-method
 (findChromPeaks-centWave), 99
noise<-,MassifquantParam-method
 (findChromPeaks-massifquant),
 109
normalize,21,22
normalize,XCMSnExp-method
 (bin,XCMSnExp-method),21
NumericList(),229</pre>

ObiwarpParam (adjustRtime-obiwarp), 7 ObiwarpParam-class (adjustRtime-obiwarp), 7 OnDiskMSnExp, 21–23, 32, 33, 45, 90, 99, 103–105, 108, 109, 114–116, 119, 120, 126, 129, 130, 182, 269, 279 overlappingFeatures, 189, 278

palette, 45 par(), 210 Param (GenericParam-class), 145 Param-class (GenericParam-class), 145 pdf. 198 PeakDensityParam, 204, 205 PeakDensityParam (groupChromPeaks-density), 155 PeakDensityParam(), 155, 258, 261 PeakDensityParam-class (groupChromPeaks-density), 155 peakDetectionCWT, 63, 64, 129, 144, 145 peakGroupsMatrix (adjustRtime-peakGroups), 13 peakGroupsMatrix, PeakGroupsParam-method (adjustRtime-peakGroups), 13 peakGroupsMatrix<-(adjustRtime-peakGroups), 13 peakGroupsMatrix<-,PeakGroupsParam-method (adjustRtime-peakGroups), 13 PeakGroupsParam, 13 PeakGroupsParam (adjustRtime-peakGroups), 13 PeakGroupsParam-class (adjustRtime-peakGroups), 13 peakPlots,xcmsSet-method (peakPlots-methods), 190 peakPlots-methods, 190 peaks (xcmsSet-class), 288

peaks,xcmsSet-method(xcmsSet-class), 288 peaks<- (xcmsSet-class), 288</pre> peaks<-,xcmsSet-method (xcmsSet-class),</pre> 288 peakScaleRange (findPeaks-MSW), 126 peakScaleRange,MSWParam-method (findPeaks-MSW), 126 peakScaleRange<- (findPeaks-MSW), 126</pre> peakScaleRange<-,MSWParam-method</pre> (findPeaks-MSW), 126 peaksWithCentWave, 104, 191, 195 peaksWithCentWave(), 96, 97 peaksWithMatchedFilter, 120, 193, 194 peaksWithMatchedFilter(), 98 peakTable (peakTable-methods), 195 peakTable,xcmsSet-method (peakTable-methods), 195 peakTable-methods, 195 peakThr (findPeaks-MSW), 126 peakThr,MSWParam-method (findPeaks-MSW), 126 peakThr<-(findPeaks-MSW), 126</pre> peakThr<-,MSWParam-method</pre> (findPeaks-MSW), 126 peakwidth (findChromPeaks-centWave), 99 peakwidth, CentWaveParam-method (findChromPeaks-centWave), 99 peakwidth, MassifquantParam-method (findChromPeaks-massifquant), 109 peakwidth<- (findChromPeaks-centWave),</pre> 99 peakwidth<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 peakwidth<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 phenoData (xcmsSet-class), 288 phenoData, xcmsSet-method (xcmsSet-class), 288 phenoData<- (xcmsSet-class), 288</pre> phenoData<-,xcmsSet,ANY-method</pre> (xcmsSet-class), 288 phenoData<-,xcmsSet-method</pre> (xcmsSet-class), 288 phenoDataFromPaths, 197 pickPeaks, 21, 22

pickPeaks,XCMSnExp-method (bin, XCMSnExp-method), 21 plot, 175, 265, 271 plot(), 258 plot, plot-methods (plot.xcmsEIC), 197 plot,XChromatogram,ANY-method (XChromatograms), 252 plot,XChromatograms,ANY-method (XChromatograms), 252 plot,XCMSnExp,missing-method (XCMSnExp-class), 269 plot.xcmsEIC, 197 plotAdjustedRtime, 7, 12, 17, 198, 278 plotChrom, 208, 221, 285 plotChrom (plotChrom-methods), 200 plotChrom, xcmsRaw-method (plotChrom-methods), 200 plotChrom-methods, 200 plotChromatogramsOverlay, 201 plotChromatogramsOverlay, MChromatograms-methopdlotSurf (plotSurf-methods), 216 (plotChromatogramsOverlay), 201 plotChromatogramsOverlay,XChromatograms-method (plotChromatogramsOverlay), 201 plotChromPeakDensity (plotChromPeakDensity,XCMSnExp-methodplotTIC (plotTIC-methods), 217 204 plotChromPeakDensity(), 158 plotChromPeakDensity,XChromatograms-method (XChromatograms), 252 plotChromPeakDensity,XCMSnExp-method, 204 plotChromPeakImage, 277 plotChromPeakImage(plotChromPeaks), 206 plotChromPeaks, 35, 206, 277 plotEIC (plotEIC-methods), 208 plotEIC, xcmsRaw-method (plotEIC-methods), 208 plotEIC-methods, 208 plotFeatureGroups, 209 plotFeatureGroups(), 74 plotMsData, 210 plotPeaks (plotPeaks-methods), 211 plotPeaks, xcmsRaw-method (plotPeaks-methods), 211 plotPeaks-methods, 211 plotQC, 212 plotRaw, 227, 285 plotRaw (plotRaw-methods), 213

plotRaw, xcmsRaw-method (plotRaw-methods), 213 plotRaw-methods, 213 plotrt, 289 plotrt (plotrt-methods), 214 plotrt,xcmsSet-method(plotrt-methods), 214plotrt-methods, 214 plotScan, 285 plotScan (plotScan-methods), 215 plotScan, xcmsRaw-method (plotScan-methods), 215 plotScan-methods, 215 plotSpec, 221, 285 plotSpec (plotSpec-methods), 215 plotSpec,xcmsRaw-method (plotSpec-methods), 215 plotSpec-methods, 215 plotSurf, 285 plotSurf,xcmsRaw-method (plotSurf-methods), 216 plotSurf-methods, 216 plotTIC, 285 plotTIC, xcmsRaw-method (plotTIC-methods), 217 plotTIC-methods, 217 plotTree (xcmsFragments-class), 268 plotTree,xcmsFragments-method (xcmsFragments-class), 268 png, 198 polarity,CentWavePredIsoParam-method (findChromPeaks-centWaveWithPredIsoROIs), 105 polarity<-(findChromPeaks-centWaveWithPredIsoROIs), 105 polarity<-.CentWavePredIsoParam-method</pre> (findChromPeaks-centWaveWithPredIsoROIs), 105 postscript, 198 ppm (findChromPeaks-centWave), 99 ppm,CentWaveParam-method (findChromPeaks-centWave), 99 ppm,FillChromPeaksParam-method (FillChromPeaksParam-class), 80 ppm,MassifquantParam-method

(findChromPeaks-massifquant), 109 ppm,MzClustParam-method (groupChromPeaks-mzClust), 159 ppm<- (findChromPeaks-centWave), 99</pre> ppm<-,CentWaveParam-method (findChromPeaks-centWave), 99 ppm<-,FillChromPeaksParam-method (FillChromPeaksParam-class), 80 ppm<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 ppm<-,MzClustParam-method (groupChromPeaks-mzClust), 159 prefilter(findChromPeaks-centWave), 99 prefilter.CentWaveParam-method (findChromPeaks-centWave), 99 prefilter, MassifquantParam-method (findChromPeaks-massifquant), 109 prefilter<- (findChromPeaks-centWave),</pre> 99 prefilter<-.CentWaveParam-method</pre> (findChromPeaks-centWave), 99 prefilter<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 present (absent-methods), 6 present, xcmsSet-method (absent-methods), 6 processDate (ProcessHistory-class), 217 processDate, ProcessHistory-method (ProcessHistory-class), 217 ProcessHistory, 258, 260, 270, 275, 277 ProcessHistory (ProcessHistory-class), 217 processHistory, 16, 145, 269 processHistory (XCMSnExp-class), 269 processHistory(), 19 processHistory, XChromatograms-method (XChromatograms), 252 processHistory,XCMSnExp-method (XCMSnExp-class), 269 ProcessHistory-class, 217 processHistoryTypes, 218 processHistoryTypes(XCMSnExp-class), 269 processInfo (ProcessHistory-class), 217

processInfo, ProcessHistory-method (ProcessHistory-class), 217 processParam (ProcessHistory-class), 217 processParam, XProcessHistory-method (ProcessHistory-class), 217 processType (ProcessHistory-class), 217 processType, ProcessHistory-method (ProcessHistory-class), 217 profBin, 221, 286, 287 profBinLin, 286 profBinLinBase, 286 profile-matrix (profMat-xcmsSet), 219 profinfo, 284, 285 profinfo (xcmsSet-class), 288 profinfo, xcmsRaw-method (xcmsRaw-class). 283 profinfo,xcmsSet-method (xcmsSet-class), 288 profinfo<- (xcmsSet-class), 288</pre> profinfo<-,xcmsSet-method</pre> (xcmsSet-class), 288 profIntLin, 286 profMat, 269, 283 profMat (profMat-xcmsSet), 219 profMat, OnDiskMSnExp-method (XCMSnExp-class), 269 profMat,XCMSnExp-method (XCMSnExp-class), 269 profMat,xcmsRaw-method (profMat-xcmsSet), 219 profMat-xcmsSet, 219 profMedFilt, 285 profMedFilt(profMedFilt-methods), 221 profMedFilt, xcmsRaw-method (profMedFilt-methods), 221 profMedFilt-methods, 221 profMethod, 221, 223, 282, 285, 287 profMethod (profMethod-methods), 221 profMethod, xcmsRaw-method (profMethod-methods), 221 profMethod, xcmsSet-method (xcmsSet-class), 288 profMethod-methods, 221 profMethod<-, 285 profMethod<- (profMethod-methods), 221</pre> profMethod<-,xcmsRaw-method</pre> (profMethod-methods), 221 profMz (xcmsRaw-class), 283

profMz,xcmsRaw-method(xcmsRaw-class), 283 profRange, 148, 149, 200, 215, 216, 285 profRange (profRange-methods), 222 profRange, xcmsRaw-method (profRange-methods), 222 profRange-methods, 222 profStep, 282, 285, 287 profStep (profStep-methods), 223 profStep,xcmsRaw-method (profStep-methods), 223 profStep,xcmsSet-method (xcmsSet-class), 288 profStep-methods, 223 profStep<-, 285 profStep<- (profStep-methods), 223</pre> profStep<-,xcmsRaw-method (profStep-methods), 223 progressCallback (xcmsSet-class), 288 progressCallback,xcmsSet-method (xcmsSet-class), 288 progressCallback<- (xcmsSet-class), 288</pre> progressCallback<-,xcmsSet-method</pre> (xcmsSet-class), 288 pSet, 279

quantify, 278
quantify, XCMSnExp-method, 224

rawEIC, 146, 147, 209 rawEIC (rawEIC-methods), 226 rawEIC,xcmsRaw-method (rawEIC-methods), 226 rawEIC-methods, 226 rawMat(rawMat-methods), 227 rawMat,xcmsRaw-method(rawMat-methods), 227 rawMat-methods, 227 reconstructChromPeakSpectra, 227 reconstructChromPeakSpectra(), 122 refineChromPeaks, 35 refineChromPeaks (CleanPeaksParam), 38 refineChromPeaks(), 96, 184 refineChromPeaks, XChromatogram, MergeNeighboringPeaksParennetheakgroups-methods), (XChromatograms), 252 refineChromPeaks, XChromatograms, MergeNeighbortegPeaksDawanpmethod/, 231, 233 (XChromatograms), 252 refineChromPeaks,XCMSnExp,CleanPeaksParam-method (CleanPeaksParam), 38 retcor.peakgroups, 17

refineChromPeaks,XCMSnExp,FilterIntensityParam-method (FilterIntensityParam), 94 refineChromPeaks,XCMSnExp,MergeNeighboringPeaksParam-metho (MergeNeighboringPeaksParam), 186 register, 103, 108, 114, 120, 129 removeIntensity (removeIntensity,Chromatogram-method), 229 removeIntensity,Chromatogram-method, 229 removeIntensity,MChromatograms-method (removeIntensity, Chromatogram-method), 229 removeIntensity,XChromatogram-method (removeIntensity,Chromatogram-method), 229 removePeaks, 21, 22 removePeaks,XCMSnExp-method (bin,XCMSnExp-method), 21 response (adjustRtime-obiwarp), 7 response,ObiwarpParam-method (adjustRtime-obiwarp), 7 response<- (adjustRtime-obiwarp), 7 response<-,ObiwarpParam-method (adjustRtime-obiwarp), 7 retcor, 7, 11, 214, 290 retcor (retcor-methods), 230 retcor,xcmsSet-method(retcor-methods), 230 retcor-methods, 230 retcor.linear (retcor.peakgroups-methods), 233 retcor.linear,xcmsSet-method (retcor.peakgroups-methods), 233 retcor.loess, 231 retcor.loess (retcor.peakgroups-methods), 233 retcor.loess,xcmsSet-method 233 retcor.obiwarp,xcmsSet-method (retcor.obiwarp), 231

retcor.peakgroups (retcor.peakgroups-methods), 233 retcor.peakgroups,xcmsSet-method (retcor.peakgroups-methods), 233 retcor.peakgroups-methods, 233 retexp, 234 revMz (xcmsRaw-class), 283 revMz,xcmsRaw-method(xcmsRaw-class), 283 ridgeLength (findPeaks-MSW), 126 ridgeLength,MSWParam-method (findPeaks-MSW), 126 ridgeLength<- (findPeaks-MSW), 126 ridgeLength<-,MSWParam-method (findPeaks-MSW), 126 rla, 234 roiList(findChromPeaks-centWave), 99 roiList,CentWaveParam-method (findChromPeaks-centWave), 99 roiList<- (findChromPeaks-centWave), 99 roiList<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 roiScales (findChromPeaks-centWave), 99 roiScales, CentWaveParam-method (findChromPeaks-centWave), 99 roiScales<- (findChromPeaks-centWave),</pre> 99 roiScales<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 rowRla(rla), 234 rtime,XCMSnExp-method(XCMSnExp-class), 269 rtrange (xcmsEIC-class), 265 rtrange,xcmsEIC-method(xcmsEIC-class), 265 sampclass, 6, 287 sampclass (xcmsSet-class), 288 sampclass,xcmsSet-method (xcmsSet-class), 288 sampclass<- (xcmsSet-class), 288</pre> sampclass<-,xcmsSet-method</pre> (xcmsSet-class), 288 sampleGroups(groupChromPeaks-density), 155 sampleGroups,MzClustParam-method

(groupChromPeaks-mzClust), 159

sampleGroups, NearestPeaksParam-method (groupChromPeaks-nearest), 162 sampleGroups,PeakDensityParam-method (groupChromPeaks-density), 155 sampleGroups<-(groupChromPeaks-density), 155 sampleGroups<-,MzClustParam-method</pre> (groupChromPeaks-mzClust), 159 sampleGroups<-,NearestPeaksParam-method</pre> (groupChromPeaks-nearest), 162 sampleGroups<-,PeakDensityParam-method</pre> (groupChromPeaks-density), 155 sampnames, 265, 290 sampnames (sampnames-methods), 235 sampnames, xcmsEIC-method (sampnames-methods), 235 sampnames, xcmsSet-method (sampnames-methods), 235 sampnames-methods, 235 sampnames<- (xcmsSet-class), 288</pre> sampnames<-,xcmsSet-method</pre> (xcmsSet-class), 288 sav.gol, 127, 129, 144 scales (findPeaks-MSW), 126 scales,MSWParam-method(findPeaks-MSW), 126 scales<- (findPeaks-MSW), 126</pre> scales<-,MSWParam-method</pre> (findPeaks-MSW), 126 scanrange(xcmsSet-class), 288 scanrange, xcmsRaw-method (xcmsRaw-class), 283 scanrange, xcmsSet-method (xcmsSet-class), 288 selfStart, 244 SerialParam. 286 setAs (XCMSnExp-class), 269 show, 268 show,CleanPeaksParam-method (CleanPeaksParam), 38 show, FilterIntensityParam-method (FilterIntensityParam), 94 show,MergeNeighboringPeaksParam-method (MergeNeighboringPeaksParam), 186 show, MsFeatureData-method (XCMSnExp-class), 269 show,ProcessHistory-method

(ProcessHistory-class), 217 show, XChromatogram-method (XChromatograms), 252 show, XChromatograms-method (XChromatograms), 252 show, xcmsEIC-method (xcmsEIC-class), 265 show, xcmsFragments-method (xcmsFragments-class), 268 show, XCMSnExp-method (XCMSnExp-class), 269 show, xcmsPeaks-method (xcmsPeaks-class), 280 show, xcmsRaw-method (xcmsRaw-class), 283 show, xcmsSet-method (xcmsSet-class), 288 show, XProcessHistory-method (ProcessHistory-class), 217 showError(showError,xcmsSet-method), 236 showError, xcmsSet-method, 236 sigma(findChromPeaks-matchedFilter), 116 sigma, MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 sigma<- (findChromPeaks-matchedFilter),</pre> 116 sigma<-.MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 SimilarRtimeParam(), 167 smooth. 21. 22 smooth(adjustRtime-peakGroups), 13 smooth,PeakGroupsParam-method (adjustRtime-peakGroups), 13 smooth,XCMSnExp-method (bin, XCMSnExp-method), 21 smooth<- (adjustRtime-peakGroups), 13</pre> smooth<-,PeakGroupsParam-method</pre> (adjustRtime-peakGroups), 13 SnowParam, 286 SnowParam(), 247 snthresh (findChromPeaks-centWave), 99 snthresh, CentWaveParam-method (findChromPeaks-centWave), 99 snthresh, MassifquantParam-method (findChromPeaks-massifquant), 109 snthresh,MatchedFilterParam-method

(findChromPeaks-matchedFilter), 116 snthresh.MSWParam-method (findPeaks-MSW), 126 snthresh<- (findChromPeaks-centWave), 99</pre> snthresh<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 snthresh<-,MassifquantParam-method</pre> (findChromPeaks-massifquant), 109 snthresh<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 snthresh<-,MSWParam-method</pre> (findPeaks-MSW), 126 snthreshIsoROIs (findChromPeaks-centWaveWithPredIsoROIs), 105 snthreshIsoROIs,CentWavePredIsoParam-method (findChromPeaks-centWaveWithPredIsoROIs), 105 snthreshIsoROIs<-</pre> (findChromPeaks-centWaveWithPredIsoROIs), 105 snthreshIsoROIs<-,CentWavePredIsoParam-method</pre> (findChromPeaks-centWaveWithPredIsoROIs), 105 sortMz (xcmsRaw-class), 283 sortMz,xcmsRaw-method(xcmsRaw-class), 283 span (adjustRtime-peakGroups), 13 span, PeakGroupsParam-method (adjustRtime-peakGroups), 13 span<- (adjustRtime-peakGroups), 13</pre> span<-,PeakGroupsParam-method</pre> (adjustRtime-peakGroups), 13 specDist(specDist-methods), 237 specDist,xcmsSet-method (specDist-methods), 237 specDist-methods, 237 specDist.cosine, 238 specDist.cosine,matrix,matrix-method (specDist.cosine), 238 specDist.meanMZmatch, 239 specDist.meanMZmatch,matrix,matrix-method (specDist.meanMZmatch), 239 specDist.peakCount (specDist.peakCount-methods),

240

specDist.peakCount,matrix,matrix-method (specDist.peakCount-methods), 240 specDist.peakCount-methods, 240 specNoise, 240, 242 specPeaks, 241, 241 spectra,XCMSnExp-method (XCMSnExp-class), 269 spectrapply,XCMSnExp-method (XCMSnExp-class), 269 Spectrum, 37, 78, 270, 277 Spectrum2, 37, 78, 229 split, 290 split, split-methods(split.xcmsSet), 243 split,XCMSnExp,ANY-method (filterFeatureDefinitions), 90 split.screen, 191, 212 split.xcmsRaw, 242, 292 split.xcmsSet, 243 SSgauss, 243 Startup, 247 steps(findChromPeaks-matchedFilter), 116 steps,MatchedFilterParam-method (findChromPeaks-matchedFilter), 116 steps<- (findChromPeaks-matchedFilter),</pre> 116 steps<-,MatchedFilterParam-method</pre> (findChromPeaks-matchedFilter), 116 stitch (stitch-methods), 244 stitch, xcmsRaw-method (stitch-methods), 244 stitch-methods, 244 stitch.netCDF (stitch-methods), 244 stitch.xml (stitch-methods), 244 structure, 281 subset(adjustRtime-peakGroups), 13 subset,ObiwarpParam-method (adjustRtime-obiwarp), 7 subset,PeakGroupsParam-method (adjustRtime-peakGroups), 13 subset-xcmsRaw 292

subset<- (adjustRtime-peakGroups), 13</pre> subset<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 subset<-,PeakGroupsParam-method</pre> (adjustRtime-peakGroups), 13 subsetAdjust(adjustRtime-peakGroups), 13 subsetAdjust,ObiwarpParam-method (adjustRtime-obiwarp), 7 subsetAdjust,PeakGroupsParam-method (adjustRtime-peakGroups), 13 subsetAdjust<-</pre> (adjustRtime-peakGroups), 13 subsetAdjust<-,ObiwarpParam-method</pre> (adjustRtime-obiwarp), 7 subsetAdjust<-,PeakGroupsParam-method</pre> (adjustRtime-peakGroups), 13 SummarizedExperiment, 224, 225, 278

adjustRtime-peakGroups), 13vector, 281ObiwarpParam-methodverboseColumns(adjustRtime-obiwarp), 7(findChromPeaks-centWave), 99ObiwarpParam-methodverboseColumns, CentWaveParam-method(adjustRtime-peakGroups), 13(findChromPeaks-centWave), 99verboseColumns, CentWaveParam-method(findChromPeaks-centWave), 99verboseColumns, MassifquantParam-method(findChromPeaks-centWave), 99verboseColumns, MassifquantParam-method(findChromPeaks-centWave), 99verboseColumns, MassifquantParam-method(findChromPeaks-massifquant), 109

verboseColumns,MSWParam-method (findPeaks-MSW), 126 verboseColumns<-(findChromPeaks-centWave), 99 verboseColumns<-,CentWaveParam-method</pre> (findChromPeaks-centWave), 99 verboseColumns<-,MassifquantParam-method (findChromPeaks-massifquant), 109 verboseColumns<-,MSWParam-method (findPeaks-MSW), 126 verify.mzQuantM, 247 verify.mzQuantML, 250 verify.mzQuantML (verify.mzQuantM), 247 withWave (findChromPeaks-massifquant), 109 withWave,MassifquantParam-method (findChromPeaks-massifquant), 109 withWave<-(findChromPeaks-massifquant), 109 withWave<-,MassifquantParam-method (findChromPeaks-massifquant), 109 write.cdf (write.cdf-methods), 248 write.cdf,xcmsRaw-method (write.cdf-methods), 248 write.cdf-methods, 248 write.mzdata(write.mzdata-methods), 249 write.mzdata,xcmsRaw-method (write.mzdata-methods), 249 write.mzdata-methods, 249 write.mzQuantML, 248 write.mzQuantML (write.mzQuantML-methods), 249 write.mzQuantML,xcmsSet-method (write.mzQuantML-methods), 249 write.mzQuantML-methods, 249 writeMSData(), 251 writeMSData,XCMSnExp,character-method, 250 writeMzTab, 251

XChromatogram, 96XChromatogram (XChromatograms), 252XChromatogram(), 230 XChromatogram-class (XChromatograms), 252 XChromatograms, 34, 252 XChromatograms(), 75, 76, 87-89, 93, 202, 230 XChromatograms-class (XChromatograms), 252 xcms-deprecated, 264 xcmsEIC-class. 265 xcmsFileSource, 290 xcmsFileSource-class, 266 xcmsFragments, 40, 267, 268, 282 xcmsFragments-class, 268 XCMSnExp, 11, 12, 16–18, 21–23, 29–34, 36, 37, 39, 45, 71, 74, 75, 77, 90, 92, 93, 95, 104, 108, 109, 115, 120, 130, 145, 157, 158, 160, 161, 164, 172, 187, 199, 205, 207, 220, 224, 225, 250, 271 XCMSnExp(XCMSnExp-class), 269 XCMSnExp(), 74, 165, 166, 168, 170, 210, 259, 261 XCMSnExp-class, 269 XCMSnExp-filter (filterFeatureDefinitions), 90 xcmsPeaks-class, 280 xcmsRaw, 40, 140-142, 144, 149, 183, 189, 219, 220, 248, 249, 269, 281, 281, 283, 285, 290-292 xcmsRaw-class, 283 xcmsSet, 40, 104, 108, 115, 120, 130, 140, 212, 236, 246, 250, 251, 268, 269, 279, 286, 288–290 xcmsSet-class, 288 xcmsSource, 184, 266, 291 xcmsSource (xcmsSource-methods), 291 xcmsSource, character-method (xcmsFileSource-class), 266 xcmsSource, xcmsSource-method (xcmsSource-methods), 291 xcmsSource-class, 290 xcmsSource-methods, 291 xdata. 291 XProcessHistory (ProcessHistory-class), 217 XProcessHistory-class (ProcessHistory-class), 217