

CAMERA

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annotate	<i>Deconvolute/Annotate LC/ESI-MS data</i>
----------	--

Description

Annotate isotope peaks, adducts and fragments for a (grouped) `xcmsSet` `xs`. Returns a `xsAnnotate` object.

Usage

```
annotate(xs, sigma = 6, perfw hm = 0.6, cor_eic_th = 0.75, maxcharge = 3, maxiso
```

Arguments

<code>xs</code>	<code>xcmsSet</code> with peak group assignments
<code>sigma</code>	Isotopic peak relationship table
<code>perfw hm</code>	Adduct/Fragment peak relationship table
<code>cor_eic_th</code>	correlation threshold (0..1)
<code>maxcharge</code>	max charge of the ions
<code>maxiso</code>	max number of isotopes
<code>ppm</code>	ppm error
<code>mzabs</code>	absolut error

multiplier	max. number n of [nM+x] clusterions
sample	Index of which sample is used for the correlation
category	Which class label should be used
polarity	Which polarity mode was used for measuring of the ms sample

Details

Batch script for a annotation for a a (grouped) xcmsSet *xs*. Generate intern a xsAnnotate object and calls the member function for the annotation step. Returns a peaklist with additional columns for isotopes, adducts and an index for the pseudo spectra group. xsAnnotate groups Peaks into spectra groups, after there Retentiontime and EIC correlations. Attention: For the EIC correlation only one sample can be used, so if the xcmsSet contains more than one sample, one must be chosen.

Value

annotate returns an xsAnnotate object. For more information about see [xsAnnotate-class](#)

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(file, method="centWave", ppm=30, peakwidth=c(5,10))
annotated_xs<- annotate(xs)
```

findAdducts-methods

Calculate Adducts and Annotate LC/ESI-MS Spectra

Description

Annotate adducts (and fragments) for a xsAnnotate object. Returns a xsAnnotate object with annotated pseudospectra.

Usage

```
findAdducts(object, ppm=5, mzabs=0.015, multiplier=3, polarity=NULL)
```

Arguments

object	the xsAnnotate object
ppm	ppm error for the search
mzabs	allowed variance for the search
multiplier	highest number(n) of allowed clusterion [nM+ion]
polarity	Which polarity mode was used for measuring of the ms sample

Details

Adducts (and fragments) are annotated for a `xsAnnotate` object. For every pseudospectra group, generated bei `groupFWHM` and `groupCorr`, all possible Adducts are calculated and mapped to the peaks. If at least two adducts match, a possible molecule-mass for the group can be calculated. After the annotation every masshypothese is checked against the charge of the calculated isotopes. It is recommend to call `findIsotopes()` before the annotation step.

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(file, method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
# an <- groupCorr(an,xs) # optional but very recommended step

an <- findIsotopes(an) # optional but recommended.
an <- findAdducts(an,polarity="positive")
peaklist <- getPeaklist(an) # get the annotated peak list
```

findIsotopes-methods

Deconvolute/Annotate LC/ESI-MS data

Description

Annotate isotope peaks for a `xsAnnotate` object. Returns a `xsAnnotate` object with annotated isotopes.

Arguments

<code>object</code>	the <code>xsAnnotate</code> object
<code>maxcharge</code>	max. number of the isotope charge
<code>maxiso</code>	max. number of the isotope peaks
<code>ppm</code>	ppm error for the search
<code>mzabs</code>	allowed variance for the search

Details

Isotope peaks are annotated for a `xsAnnotate` object according to given rules (`maxcharge`, `maxiso`). The algorithm benefits from a earlier grouping of the data, with `groupFWHM` and `groupCorr`. Generates a list of all possible isotopes, which is stored in `object@isotopes`.

Methods

```
object = "xsAnnotate" findIsotopes(object, maxcharge=3, maxiso=3, ppm=5, mzabs=0.01)
```

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(file, method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
an <- findIsotopes(an)
```

getPeaklist

Retrieve the annotated peaklist

Description

Extract all groups from a xsAnnotate object. Returns a peaklist with annotated peaks.

Usage

```
getPeaklist(object)
```

Arguments

object xsAnnotate object

Details

xsAnnotate groups LC/MS Peaklist after there EIC correlation and FWHM. These function extract one of these so called "pseudo spectra groups" with include the peaklist with there annotations.

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(c(file), method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
an <- findIsotopes(an)
an <- findAdducts(an,polarity="positive")
peaklist <- getPeaklist(an)
```

getpspectra *Retrieve a spectra-group peaklist*

Description

Extract one group from a xsAnnotate object. Returns a peaklist with annotated peaks.

Usage

```
getpspectra(object, grp)
```

Arguments

object	xsAnnotate object
grp	index of pseudo-spectra-group

Details

xsAnnotate groups LC/MS Peaklist after there EIC correlation and FWHM. These function extract one of these so called "pseudo spectra groups" with include the peaklist with there annotations. The annotation depends on a before called findAdducts() (and findIsotopes()). Attention: The indices for the isotopes, are those from the whole peaklist. See getPeaklist() .

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(c(file), method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
peaklist <- getpspectra(an, 1)
```

groupCorr-methods *EIC correlation grouping of LC/ESI-MS data*

Description

Grouping the peaks after the correlation of the EICs into pseudospectragroups for a xsAnnotate object. Return a xsAnnotate object with group information.

Usage

```
groupCorr(object, cor_eic_th=0.75)
```

Arguments

object the xsAnnotate object
cor_eic_th correlation threshold for the EIC correlation

Details

The algorithm correlates the EIC of a every peak with all others, to find the peaks that belong to one substance. LC/MS data should grouped with groupFWHM first. This step reduce the runtime a lot and increased the number of correct classifications.

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(c(file), method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
an <- groupCorr(an)
```

groupFWHM-methods *FWHM-Grouping of LC/ESI-MS data*

Description

Grouping the peaks after the FWHM of the retentiontimes into pseudospectragroups for a xsAnnotate object. Returns a xsAnnotate object with group information.

Usage

```
groupFWHM(object, sigma = 6 , perfwhm = 0.6)
```

Arguments

object the xsAnnotate object
sigma the multiplier of the standard deviation
perfwhm percentage of the width of the FWHM

Details

The FWHM (full width at half maximum) of a peak, will be used for the grouping. Every peak who eluated at the same time as a selected peak, will be part of the group. Same time is defined about the $Rt-med \pm FWHM * perfwhm$. FWHM is calculated as the product of $\sigma * standard\ deviation$.

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Examples

```
library(CAMERA)
file <- system.file('mzdata/MM14.mzdata', package = "CAMERA")
xs <- xcmsSet(file, method="centWave", ppm=30, peakwidth=c(5,10))
an <- xsAnnotate(xs)
an <- groupFWHM(an)
```

mm14

*Extract of marker mixture 14 LC/MS data***Description**

xcmsSet object containing quantitated LC/MS peaks from a marker mixture. The data is a centroided subset from 117-650 m/z and 271-302 seconds with 134 peaks. Positive ionization mode data in mzData file format.

Usage

```
data(mm14)
```

Format

The format is:

```
Formal class 'xcmsSet' [package "xcms"] with 8 slots
  @ peaks      : num [1:83, 1:11] 117 117 118 119 136
  .. ..- attr(*, "dimnames")=List of 2
  .. .. ..$ : NULL
  .. .. ..$ : chr [1:11] "mz" "mzmin" "mzmax" "rt"
  ..@ groups   : logi[0 , 0 ]
  ..@ groupidx : list()
  ..@ phenoData:'data.frame':  1 obs. of  1 variable:
  .. ..$ class: Factor w/ 1 level "mzdata": 1
  ..@ rt       :List of 2
  .. ..$ raw    :List of 1
  .. .. ..$ : num [1:112] 270 271 271 271 272 ...
  .. ..$ corrected:List of 1
  .. .. ..$ : num [1:112] 270 271 271 271 272 ...
  ..@ filepaths: chr "mzdata/MM14.mzdata"
  ..@ profinfo :List of 2
  .. ..$ method: chr "bin"
  .. ..$ step   : num 0.1
  ..@ polarity  : chr(0)
```

Details

The corresponding raw mzData files are located in the mzData subdirectory of this package.

Author(s)

Carsten Kuhl <ckuhl@ipb-halle.de>

Source

<http://doi:10.1186/1471-2105-9-504>

References

Data originally reported in "Highly sensitive feature detection for high resolution LC/MS" BMC Bioinformatics; 2008; 9:504.

xsAnnotate-class *Class xsAnnotate, a class for annotate peak data*

Description

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

Objects from the Class

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

Slots

peaks: matrix containing peak data
pspectra: list containing peak data
isotopes: matrix containing peak data
derivativeIons: matrix containing peak data
formula: matrix containing peak data
sample: the number of the used xcmsSet sample

Methods

groupFWHM signature(object = "xsAnnotate"): group the peak data after the FWHM of the retention time
groupCorr signature(object = "xsAnnotate"): group the peak data after the Correlation of the EIC
findIsotopes signature(object = "xsAnnotate"): search for possible isotopes in the spectra
findAdducts signature(object = "xsAnnotate"): search for possible adducts in the spectra

Note

No notes yet.

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See Also

[xsAnnotate](#)

xsAnnotate	<i>Constructor for a xsAnnotate object which one assigned xcmsSet object</i>
------------	--

Description

This function handles the construction of xsAnnotate object. It is generate from a xcmsSet object adapt its peaktable.

Usage

```
xsAnnotate(xs = NULL, sample=NA, category=NA)
```

Arguments

xs	a xcmsSet object
sample	Index of the group xcmsSet sample, that is used for the EIC corelations step.
category	Class label, for the class label of a grouped xcmsSet with different classes.

Value

A xsAnnotate object.

Author(s)

Carsten Kuhl, <ckuhl@ipb-halle.de>

See Also

[xsAnnotate-class](#)

Examples

```
library(faahKO)
xs <- group(faahko)
xs_anno <- xsAnnotate(xs, sample=1, category="WT")
```

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