

Package ‘CluMSID’

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Type Package

Title Clustering of MS2 Spectra for Metabolite Identification

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Description CluMSID is a tool that aids the identification of features in untargeted LC-MS/MS analysis by the use of MS2 spectra similarity and unsupervised statistical methods. It offers functions for a complete and customisable workflow from raw data to visualisations and is interfaceable with the xcms family of preprocessing packages.

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URL <https://github.com/tdepke/CluMSID>

BugReports <https://github.com/tdepke/CluMSID/issues>

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R topics documented:

accessors	2
addAnnotations	4
as.MS2spectrum	5
cossim	5
distanceMatrix	6
extractMS2spectra	7
extractPseudospectra	8
featureList	9
findFragment	10
findNL	10
getSimilarities	11
getSpectrum	12
HCplot	13
HCtbl	14
MDSplot	14
mergeMS2spectra	15
MS2spectrum-class	16
networkplot	18
OPTICSplot	19
OPTICStbl	20
pseudospectrum-class	21
specplot	21
splitPolarities	22
writeFeaturelist	22
Index	24

accessors	<i>Accessor functions for individual slots of MS2spectrum and pseudospectrum objects</i>
-----------	--

Description

Accessor functions for individual slots of [MS2spectrum](#) and [pseudospectrum](#) objects

Usage

```
accessID(x)

accessAnnotation(x)

accessPrecursor(x)

accessRT(x)

accessPolarity(x)
```

```
accessSpectrum(x)
```

```
accessNeutralLosses(x)
```

Arguments

x An object of class `MS2spectrum` or `pseudospectrum`

Value

The value of the respective slot of the object (id, annotation, precursor, rt, spectrum, neutral_losses)

Examples

```
load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessID(annotatedSpecList[[1]])

load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessAnnotation(annotatedSpecList[[1]])

load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessPrecursor(annotatedSpecList[[1]])

load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessRT(annotatedSpecList[[1]])

load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessPolarity(annotatedSpecList[[1]])

load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

accessSpectrum(annotatedSpecList[[1]])

load(file = system.file("extdata",
```

```

      "annotatedSpecList.RData",
      package = "CluMSIDdata"))

accessNeutralLosses(annotatedSpecList[[1]])

```

addAnnotations	<i>Adding external annotations to list of MS2spectrum objects</i>
----------------	---

Description

addAnnotations is used to add annotations that have been assigned externally, e.g. by library search, to a list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList.

Usage

```
addAnnotations(featlist, annolist, annotationColumn = 4)
```

Arguments

featlist	A list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList
annolist	A list of annotations, either as a data.frame or csv file. The order of features must be the same as in featlist. Please see the package vignette for a detailed example!
annotationColumn	The column of annolist where the annotation is found. Default is 4, which is the case if writeFeaturelist followed by manual addition of annotations, e.g. in Excel, is used to generate annolist.

Value

A list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList with external annotations added to the annotation slot of each MS2spectrum object.

Examples

```

load(file = system.file("extdata",
  "featlist.RData",
  package = "CluMSIDdata"))

addAnnotations(featlist, system.file("extdata",
  "post_anno.csv",
  package = "CluMSIDdata"),
  annotationColumn = 4)

```

as.MS2spectrum	<i>Convert spectra from MSnbase classes</i>
----------------	---

Description

Convert spectra from **MSnbase** classes

Usage

```
as.MS2spectrum(x)
```

Arguments

x An object of class [Spectrum](#) or [Spectrum2](#)

Value

An object of class [MS2spectrum](#)

Examples

```
#Load a "Spectrum2" object from MSnbase
library(MSnbase)
sp <- itraqdata[["X1"]]
#Convert this object to "MS2spectrum" class
new_sp <- as.MS2spectrum(sp)
#Or alternatively:
new_sp <- as(sp, "MS2spectrum")
```

cossim	<i>Calculate cosine similarity between two spectra</i>
--------	--

Description

cossim() calculates the cosine of the spectral contrast angle as a measure for the similarity of two spectra.

Usage

```
cossim(x, y, type = c("spectrum", "neutral_losses"),
      mzTolerance = 1e-05)

## S4 method for signature 'MS2spectrum,MS2spectrum'
cossim(x, y, type = c("spectrum",
  "neutral_losses"), mzTolerance = 1e-05)
```

```
## S4 method for signature 'pseudospectrum,pseudospectrum'
cossim(x, y,
       type = c("spectrum", "neutral_losses"), mzTolerance = 1e-05)
```

Arguments

x, y	MS2 spectra, either as matrix, MS2spectrum or pseudospectrum objects. x and y must have the same class.
type	Whether similarity between spectra ("spectrum", default) or neutral loss patterns ("neutral_losses") is to be compared
mzTolerance	The m/z tolerance used for merging. If two fragment peaks are within tolerance, they are regarded as the same. Defaults to 1e-5, i.e. 10ppm.

Value

The cosine similarity of x and y

Methods (by class)

- x = MS2spectrum, y = MS2spectrum: cossim method for [MS2spectrum](#) objects
- x = pseudospectrum, y = pseudospectrum: cossim method for [pseudospectrum](#) objects

Examples

```
load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))

cossim(annotatedSpecList[[1]], annotatedSpecList[[2]])
```

distanceMatrix	<i>Create distance matrix from list of spectra</i>
----------------	--

Description

distanceMatrix() creates a distance matrix from a list of MS2 spectra, MS1 pseudospectra or neutral loss patterns by pairwise comparison using the specified distance function. This distance matrix is the basis for CluMSID's data mining functions.

Usage

```
distanceMatrix(specList, distFun = "cossim", type = c("spectrum",
  "neutral_losses"), mz_tolerance = 1e-05)
```

Arguments

speclist	A list of MS2spectrum or pseudospectrum objects as generated by extractMS2spectra or extractPseudospectra .
distFun	The distance function to be used. At the moment, only cossim is implemented.
type	"spectrum" (default) for MS2 spectra or MS1 pseudospectra or "neutral_losses" for neutral loss patterns.
mz_tolerance	The m/z tolerance to be used for merging, default is 1e-5, i.e. +/- 10ppm. If the mass-to-charge ratios of two peaks differ less than <i>mz_tolerance</i> , they are assumed to have the same m/z .

Value

A numeric `length(speclist)` by `length(speclist)` matrix containing pairwise distances (1 - similarity) between all features in `speclist`. Row and column names are taken from the `id` slot or, if present, pasted from the `id` and `annotation` slots of the [MS2spectrum](#) or [pseudospectrum](#) objects.

Examples

```
load(file = system.file("extdata",
  "annotatedSpeclist.RData",
  package = "CluMSIDdata"))

distanceMatrix(annotatedSpeclist[1:20])
```

extractMS2spectra	<i>Extract MS2 spectra from raw data files</i>
-------------------	--

Description

`extractMS2spectra()` is used to extract MS2 spectra from raw data files, e.g. `mzXML` files.

Usage

```
extractMS2spectra(MSfile, min_peaks = 2, recalibrate_precursor = FALSE,
  RTlims = NULL)
```

Arguments

MSfile	An LC-MS/MS raw data file in one of the non-proprietary formats that can be parsed by <code>mzR</code> , e.g. <code>mzXML</code> or <code>mzML</code> .
min_peaks	Minimum number of peaks in MS2 spectrum, defaults to 2. Spectra with less than <code>min_peaks</code> fragment peaks will be ignored and not extracted.

recalibrate_precursor	Logical, defaults to FALSE. Applicable only for files that were exported to mzXML using a deprecated version of Bruker Compass Xport (< 3.0.13). If set to TRUE, the precursor m/z will be recalculated from the respective fragment m/z in the MS2 spectrum. For details, see Depke et al. 2017.
RTlims	Retention time interval for the extraction of spectra. Provide as numeric vector of length 2. Spectra with retention time < RTlims[1] or > RTlims[2] will be ignored.

Value

A list with objects of class MS2spectrum, containing MS2 spectra extracted from the raw data.

Examples

```
my_spectra <- extractMS2spectra(MSfile = system.file("extdata",
  "PoolA_R_SE.mzXML",
  package = "CluMSIDdata"),
  min_peaks = 4, RTlims = c(0,10))
```

extractPseudospectra *Extract pseudospectra*

Description

extractPseudospectra() is used to extract MS1 pseudospectra from **CAMERA** output.

Usage

```
extractPseudospectra(x, min_peaks = 1, intensity_columns = NULL)
```

Arguments

x	CAMERA output that contains information on pseudospectra. Can either be of class data.frame or xsAnnotate . It is recommended to use either xsAnnotate objects or data.frames generated from XCMSOnline results tables but other data.frames are possible.
min_peaks	Minimum number of peaks in pseudospectrum, defaults to 1. See extractMS2spectra .
intensity_columns	Numeric, defaults to NULL. If a data.frame is used as input which has not been generated from an XCMSOnline results table, the indices of the columns that contain the peak intensities in the different samples have to be indicated as intensity_columns.

Value

A list of pseudospectra, stored as objects of class [pseudospectrum](#), analogous to the output of [extractMS2spectra](#).

Examples

```
pstable <- readr::read_delim(file = system.file("extdata",
  "TD035_XCMS.annotated.diffreport.tsv",
  package = "CluMSIDdata"), delim = "\t")

pseudospeclist <- extractPseudospectra(pstable, min_peaks = 2)
```

featureList	<i>Generate a data.frame with feature information from list of MS2spectrum objects</i>
-------------	--

Description

featureList generates a data.frame that contains feature ID, precursor m/z and retention time for all features contained in a list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList. featureList is used internally by [writeFeaturelist](#).

Usage

```
featureList(featlist)
```

Arguments

featlist	A list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList
----------	--

Details

Although originally designed for lists of MS2spectrum objects, the function also works with lists of pseudospectrum objects. In this case, NA is given for precursor m/z .

Value

A data.frame that contains feature ID, precursor m/z (if available) and retention time

Examples

```
load(file = system.file("extdata",
  "featlist.RData",
  package = "CluMSIDdata"))

pre_anno <- featureList(featlist)
```

findFragment	<i>Find spectra that contain a specific fragment</i>
--------------	--

Description

findFragment is used to find spectra that contain a specific fragment ion. Its sister function is [findNL](#), which finds specific neutral losses. Both functions work analogous to [getSpectrum](#).

Usage

```
findFragment(featlist, mz, tolerance = 1e-05)
```

Arguments

featlist	a list that contains only objects of class MS2spectrum
mz	The mass-to-charge ratio of the fragment ion of interest.
tolerance	The m/z tolerance for the fragment ion search. Default is $1\text{E-}05$, i.e. +/- 10ppm.

Value

If the respective fragment is only found in one spectrum, the output is an object of class [MS2spectrum](#); if it is found in more than one spectrum, the output is a list of [MS2spectrum](#) objects.

Examples

```
load(file = system.file("extdata",
  "annotatedSpeclist.RData",
  package = "CluMSIDdata"))
putativeAQs <- findFragment(annotatedSpeclist, 159.068)
```

findNL	<i>Find spectra that contain a specific neutral loss</i>
--------	--

Description

findNL is used to find spectra that contain a specific neutral loss. Its sister function is [findFragment](#), which finds specific fragment ions. Both functions work analogous to [getSpectrum](#).

Usage

```
findNL(featlist, mz, tolerance = 1e-05)
```

Arguments

featlist	a list that contains only objects of class MS2spectrum
mz	The mass-to-charge ratio of the neutral loss of interest.
tolerance	The m/z tolerance for the neutral loss search. Default is 1E-05, i.e. +/- 10ppm.

Value

If the respective neutral loss is only found in one spectrum, the output is an object of class [MS2spectrum](#); if it is found in more than one spectrum, the output is a list of [MS2spectrum](#) objects.

Examples

```
load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))
findNL(annotatedSpecList, 212.009)
```

getSimilarities	<i>Match one spectrum against a set of spectra</i>
-----------------	--

Description

getSimilarities calculates the similarities of one spectrum or neutral loss pattern to a set of other spectra or neutral loss patterns.

Usage

```
getSimilarities(spec, speclist, type = c("spectrum", "neutral_losses"),
  hits_only = FALSE)
```

Arguments

spec	The spectrum to be compared to other spectra. Can be either an object of class MS2spectrum or a two-column numerical matrix that contains fragment mass-to-charge ratios in the first and intensities in the second column.
speclist	The set of spectra to which spec is to be compared. Must be a list where every entry is an object of class MS2spectrum . Can be generated from an mzXML file with extractMS2spectra and mergeMS2spectra or constructed using <code>new("MS2spectrum", ...)</code> for every list entry (see vignette for details).
type	Specifies whether MS2 spectra or neutral loss patterns are to be compared. Must be either 'spectrum' (default) or 'neutral_losses'.
hits_only	Logical that indicates whether the result should contain only similarities greater than zero.

Value

A named vector with similarities of spec to all spectra or neutral loss patterns in speclist.

Examples

```
load(file = system.file("extdata",
  "annotatedSpecList.RData",
  package = "CluMSIDdata"))
getSimilarities(annotatedSpecList[[137]],
  annotatedSpecList, hits_only = TRUE)
```

getSpectrum	<i>Access individual spectra from a list of spectra by various slot entries</i>
-------------	---

Description

As accessing S4 objects within lists is not trivial, getSpectrum can be used to access individual or several [MS2spectrum](#) objects by their slot entries.

Usage

```
getSpectrum(featlist, slot, what, mz.tol = 1e-05, rt.tol = 30)
```

Arguments

featlist	a list that contains only objects of class MS2spectrum
slot	The slot to be searched (invalid slot arguments will produce errors). Possible values are: • 'id' • 'annotation' • 'precursor' (<i>m/z</i> of precursor ion) • 'rt' (retention time of precursor)
what	the search term or number, must be character for 'id' and 'annotation' and numeric for 'precursor' and 'rt' See vignette for examples.
mz.tol	the tolerance used for precursor ion <i>*m/z*</i> searches, defaults to 1E-05 (+/- 10ppm)
rt.tol	the tolerance used for precursor ion retention time searches, defaults to 30s; high values can be used to specify retention time ranges (see vignette for example)

Value

If the only one spectrum matches the search criteria, the output is an object of class [MS2spectrum](#); if more than one spectrum matches, the output is a list of [MS2spectrum](#) objects.

Examples

```
load(file = system.file("extdata",
  "annotatedSpeclist.RData",
  package = "CluMSIDdata"))

getSpectrum(annotatedSpeclist, "annotation", "pyocyanin")

getSpectrum(annotatedSpeclist, "id", "M244.17T796.4")

getSpectrum(annotatedSpeclist, "precursor", 286.18, mz.tol = 1E-03)

six_eight <- getSpectrum(annotatedSpeclist, "rt", 420, rt.tol = 60)
```

HCplot	<i>Generate cluster dendrogram or heatmap from spectral similarity data</i>
--------	---

Description

HCplot() performs hierarchical clustering of spectral similarity data using average linkage as agglomeration criterion like [Hctbl](#) and generates either a circular dendrogram or a combination of dendrogram and heatmap.

Usage

```
HCplot(distmat, h = 0.95, type = c("dendrogram", "heatmap"), ...)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
h	Height where the tree is to be cut, defaults to 0.95. See cutree for details.
type	Specifies which visualisation is to be generated: "dendrogram" (default) for a circular dendrogram or "heatmap" for a combination of dendrogram and heatmap.
...	Additional graphical parameters passed to <code>plot.phylo</code> (for type = "dendrogram") or <code>gplots::heatmap.2</code> (for type = "heatmap")

Value

A plot as specified by type.

Examples

```
load(file = system.file("extdata",
  "distmat.RData",
  package = "CluMSIDdata"))

HCplot(distmat[1:50,1:50], h = 0.8, type = "heatmap")
```

H Ctbl	<i>Hierarchical clustering of spectral similarity data</i>
--------	--

Description

H Ctbl() performs hierarchical clustering of spectral similarity data using average linkage as agglomeration criterion.

Usage

```
H Ctbl(distmat, h = 0.95)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
h	Height where the tree is to be cut, defaults to 0.95. See cutree for details.

Value

A data.frame with name and cluster ID for each feature in distmat.

See Also

[HCplot](#)

Examples

```
load(file = system.file("extdata",
  "distmat.RData",
  package = "CluMSIDdata"))

my_HCtbl <- H Ctbl(distmat[1:50,1:50], h = 0.8)
```

MDSplot	<i>Multidimensional scaling of spectral similarity data</i>
---------	---

Description

MDSplot() is used to generate multidimensional scaling plots from spectral similarity data. An interactive visualisation can be produced using [plotly](#).

Usage

```
MDSplot(distmat, interactive = FALSE, highlight_annotated = FALSE, ...)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
interactive	Logical, defaults to FALSE. If TRUE, an interactive visualisation is generated using plotly .
highlight_annotated	Logical, defaults to FALSE. If TRUE, points for features for which an annotation was added before using distanceMatrix are highlighted by red colour, while other points are grey in the MDS plot.
...	Additional arguments passed to <code>geom_point()</code> , e.g. <code>pch</code> , <code>size</code> or <code>alpha</code> .

Value

An MDS plot generated with the help of [cmdscale](#), [ggplot](#) and, if interactive, [ggplotly](#).

Examples

```
load(file = system.file("extdata",
  "distmat.RData",
  package = "CluMSIDdata"))

MDSplot(distmat, highlight_annotated = TRUE)
```

mergeMS2spectra	<i>Merge MS2 spectra with or without external peak table</i>
-----------------	--

Description

mergeMS2spectra is used to merge MS2 spectra that come from the same precursor. It does so either by grouping spectra of the same precursor m/z that fall into a defined retention time window (`rt_tolerance`) or by grouping spectra to peaks from an externally supplied peak table. Please see the vignette for more details.

Usage

```
mergeMS2spectra(ms2list, mz_tolerance = 1e-05, rt_tolerance = 30,
  peaktable = NULL, exclude_unmatched = FALSE)
```

Arguments

ms2list	A list of MS2spectrum objects to be merged.
mz_tolerance	The m/z tolerance to be used for merging, default is $1e-5$, i.e. ± 10 ppm. If the mass-to-charge ratios of two peaks differ less than <code>mz_tolerance</code> , they are assumed to have the same m/z .

rt_tolerance	The retention time tolerance used for merging features. If used without a peak table, rt_tolerance is the maximum retention time difference between to subsequent spectra of the same precursor m/z with which they are still assumed to belong to the same feature. If used with an external peak table, rt_tolerance is the maximum retention time difference between a spectrum and a peak in the peak table with which the spectrum is still considered to belong to that peak.
peaktable	An external peak table, e.g. from XCMS, that serves as a template for grouping spectra. The peaktable must be a three-column data.frame with feature ID, m/z and retention time for each peak/feature.
exclude_unmatched	If an external peak table is used: Should spectra that do not match to any peak/feature in the peak table be exclude from the resulting list?

Value

A merged list of `MS2spectrum` objects.

Examples

```
my_spectra <- extractMS2spectra(MSfile = system.file("extdata",
  "Pool1A_R_SE.mzXML",
  package = "CluMSIDdata"),
  min_peaks = 4, RTlims = c(0,5))

my_merged_spectra <- mergeMS2spectra(my_spectra, rt_tolerance = 20)
```

MS2spectrum-class	<i>A custom S4 class for MS2 spectra, neutral loss patterns and respective metainformation</i>
-------------------	--

Description

A custom S4 class for MS2 spectra, neutral loss patterns and respective metainformation

Usage

```
## S4 method for signature 'MS2spectrum'
show(object)

## S4 method for signature 'MS2spectrum'
precursorMz(object)

## S4 method for signature 'MS2spectrum'
rtime(object)

## S4 method for signature 'MS2spectrum'
```



```

intensity(object)

## S4 method for signature 'MS2spectrum'
mz(object)

## S4 method for signature 'MS2spectrum,ANY'
peaksCount(object)

```

Arguments

object An object of class `MS2spectrum`

Value

Prints information from the object slots with exception of 'spectrum' and 'neutral_losses' where only a summary is given.

Methods (by generic)

- show: A show generic for MS2spectra.
- precursorMz: Method forMSnbase::precursorMz for `MS2spectrum` objects. Accesses precursor slot and returns precursor m/z as a numeric.
- rttime: Method forMSnbase::rttime for `MS2spectrum` objects. Accesses rt slot and returns retention time as a numeric.
- intensity: Method forMSnbase::intensity for `MS2spectrum` objects. Accesses spectrum slot and returns the intensity column as a numeric vector.
- mz: Method forMSnbase::mz for `MS2spectrum` objects. Accesses spectrum slot and returns the m/z column as a numeric vector.
- peaksCount: Method forMSnbase::mz for `MS2spectrum` objects. Accesses spectrum slot and returns the number of peaks as a numeric.

Slots

id a character string similar to the ID used by XCMSOnline or the ID given in a predefined peak list

annotation a character string containing a user-defined annotation, defaults to empty

precursor (median) m/z of the spectrum's precursor ion

rt (median) retention time of the spectrum's precursor ion

polarity the ionisation polarity, "positive" or "negative"

spectrum the actual MS2 spectrum as two-column matrix (column 1 is (median) m/z , column 2 is (median) intensity of the product ions)

neutral_losses a neutral loss pattern generated by subtracting the product ion mass-to-charge ratios from the precursor m/z in a matrix format analogous to the spectrum slot

networkplot

*Correlation network from spectral similarity data***Description**

networkplot() is used to generate correlation networks from spectral similarity data. An interactive visualisation can be produced using **plotly**.

Usage

```
networkplot(distmat, interactive = FALSE, show_labels = FALSE,
            label_size = 1.5, highlight_annotated = FALSE,
            min_similarity = 0.1, exclude_singletons = FALSE)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
interactive	Logical, defaults to FALSE. If TRUE, an interactive visualisation is generated using plotly .
show_labels	Logical, defaults to FALSE. If TRUE, feature IDs are printed as labels in the network plot. Argument has no effect if interactive is TRUE (because in this case, labels are displayed on mouse-over).
label_size	Numeric, defaults to 1.5. If show_labels is TRUE and interactive is FALSE, label_size defines the size of labels in the plot.
highlight_annotated	Logical, defaults to FALSE. If TRUE, points for features for which an annotation was added before using distanceMatrix are highlighted by red colour, while other points are grey in the network plot.
min_similarity	Numeric, defaults to 0.1. The minimum spectral contrast angle (see cossim) that is considered a spectral similarity and hence a connection in the network.
exclude_singletons	Logical, defaults to FALSE. If TRUE, features that have no connection to any other feature will not be displayed in the network plot.

Value

A network plot generated with the help of [network](#), [ggnet2](#) and, if interactive, [ggplotly](#). Edge weights correspond to spectral similarities.

Examples

```
load(file = system.file("extdata",
                        "distmat.RData",
                        package = "CluMSIDdata"))

networkplot(distmat[1:50,1:50], show_labels = TRUE,
```

```
exclude_singletons = TRUE)
```

OPTICSplot

Visualisation of density-based clustering of spectral similarity data

Description

OPTICSplot() performs density-based clustering of spectral similarity data using the OPTICS algorithm like [OPTICSstbl](#) and creates a reachability distance plot.

Usage

```
OPTICSplot(distmat, eps = 10000, minPts = 3, eps_cl = 0.5, ...)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
eps	OPTICS parameters, see optics .
minPts	OPTICS parameters, see optics .
eps_cl	The reachability distance used for cluster determination, see extractDBSCAN .
...	Additional graphical parameters to be passed to plot()

Details

The function internally uses [optics](#) and [extractDBSCAN](#) from the **dbscan** package.

Value

A reachability distance plot as visualisation of OPTICS clustering, see code [extractDBSCAN](#).

See Also

[OPTICSstbl](#)

Examples

```
load(file = system.file("extdata",
  "distmat.RData",
  package = "CluMSIDdata"))

OPTICSplot(distmat[1:50,1:50], eps_cl = 0.7)
```

OPTICStbl

*Density-based clustering of spectral similarity data***Description**

OPTICStbl() performs density-based clustering of spectral similarity data using the OPTICS algorithm.

Usage

```
OPTICStbl(distmat, eps = 10000, minPts = 3, eps_cl = 0.5)
```

Arguments

distmat	A distance matrix as generated by distanceMatrix .
eps, minPts	OPTICS parameters, see optics .
eps_cl	The reachability distance used for cluster determination, see extractDBSCAN .

Details

The function internally uses [optics](#) and [extractDBSCAN](#) from the **dbscan** package.

Value

A data.frame with feature name, cluster ID and OPTICS order for each feature in distmat.

See Also

[OPTICSplot](#)

Examples

```
load(file = system.file("extdata",
  "distmat.RData",
  package = "CluMSIDdata"))

my_OTPICStbl <- OPTICStbl(distmat[1:50,1:50], eps_cl = 0.7)
```

pseudospectrum-class	<i>A custom S4 class for MS1 pseudospectra and respective metainformation</i>
----------------------	---

Description

A custom S4 class for MS1 pseudospectra and respective metainformation

Slots

`id` a the "pcgroup" number assigned by **CAMERA**
`annotation` a character string containing a user-defined annotation, defaults to empty
`rt` (median) retention time of the ions contained in the pseudospectrum
`spectrum` the actual MS1 pseudospectrum as two-column matrix (column 1 is (median) m/z , column 2 is (median) intensity of the ions)

specplot	<i>Create a basic plot of MS2 spectra</i>
----------	---

Description

specplot creates a very basic plot of MS2 spectra from [MS2spectrum](#) or [pseudospectrum](#) objects.

Usage

```
specplot(spec, ...)
```

Arguments

<code>spec</code>	An object of class MS2spectrum or pseudospectrum
<code>...</code>	Additional graphical parameters to be passed to <code>plot()</code>

Value

A plot of the MS2 spectrum saved in the `spectrum` slot of `spec`.

Examples

```
load(file = system.file("extdata",  
  "annotatedSpecList.RData",  
  package = "CluMSIDdata"))  
  
specplot(annotatedSpecList[[1]])
```

splitPolarities	<i>Separate spectra with different polarities from the same run</i>
-----------------	---

Description

Using `splitPolarities`, spectra with different polarities from the same run can be separated, e.g. when processing spectra recorded with polarity-switching.

Usage

```
splitPolarities(ms2list, polarity = c("positive", "negative"))
```

Arguments

<code>ms2list</code>	A list of MS2spectrum objects as produced by extractMS2spectra .
<code>polarity</code>	The polarity of spectra to be analysed, must be "positive" or "negative".

Value

A list of [MS2spectrum](#) objects that contains only spectra with the given polarity.

Examples

```
my_spectra <- extractMS2spectra(MSfile = system.file("extdata",
  "PoolA_R_SE.mzXML",
  package = "CluMSIDdata"),
  min_peaks = 4, RTlims = c(0,5))

my_positive_spectra <- splitPolarities(my_spectra, "positive")
```

writeFeaturelist	<i>Write feature information from list of MS2spectrum objects</i>
------------------	---

Description

`writeFeaturelist` uses [featureList](#) to generate a data.frame that contains feature ID, precursor m/z and retention time for all features contained in a list of [MS2spectrum](#) objects as produced by `extractMS2spectra` and `mergeSpecList` and writes it to a csv file.

Usage

```
writeFeaturelist(featlist, filename = "pre_anno.csv")
```

Arguments

featlist	A list of MS2spectrum objects as produced by extractMS2spectra and mergeSpecList
filename	The desired file name of the csv file, default is "pre_anno.csv"

Details

Although originally designed for lists of MS2spectrum objects, the function also works with lists of pseudospectrum objects. In this case, NA is given for precursor m/z .

Value

A csv file that contains feature ID, precursor m/z and retention time. The file has a header but no row names and is separated by ' , '.

Examples

```
load(file = system.file("extdata",
  "featlist.RData",
  package = "CluMSIDdata"))

writeFeaturelist(featlist, filename = "pre_anno.csv")
```

Index

accessAnnotation (accessors), 2
accessID (accessors), 2
accessNeutralLosses (accessors), 2
accessors, 2
accessPolarity (accessors), 2
accessPrecursor (accessors), 2
accessRT (accessors), 2
accessSpectrum (accessors), 2
addAnnotations, 4
as.MS2spectrum, 5

cmdscale, 15
cossim, 5, 7, 18
cossim, MS2spectrum, MS2spectrum-method
 (cossim), 5
cossim, pseudospectrum, pseudospectrum-method
 (cossim), 5
cutree, 13, 14

distanceMatrix, 6, 13–15, 18–20

extractDBSCAN, 19, 20
extractMS2spectra, 7, 7, 8, 11, 22
extractPseudospectra, 7, 8

featureList, 9, 22
findFragment, 10, 10
findNL, 10, 10

getSimilarities, 11
getSpectrum, 10, 12
ggnet2, 18
ggplot, 15
ggplotly, 15, 18

HCplot, 13, 14
HCtbl, 13, 14

intensity, MS2spectrum-method
 (MS2spectrum-class), 16

MDSplot, 14
mergeMS2spectra, 11, 15
MS2spectrum, 2, 3, 5–7, 10–12, 16, 17, 21, 22
MS2spectrum-class, 16
mz, MS2spectrum-method
 (MS2spectrum-class), 16

network, 18
networkplot, 18

optics, 19, 20
OPTICSplot, 19
OPTICStbl, 19, 20

peaksCount, MS2spectrum, ANY-method
 (MS2spectrum-class), 16
precursorMz, MS2spectrum-method
 (MS2spectrum-class), 16
pseudospectrum, 2, 3, 6–8, 21
pseudospectrum-class, 21

rtime, MS2spectrum-method
 (MS2spectrum-class), 16

show, MS2spectrum-method
 (MS2spectrum-class), 16
specplot, 21
Spectrum, 5
Spectrum2, 5
splitPolarities, 22

writeFeaturelist, 4, 9, 22

xsAnnotate, 8