

Package ‘MetaboList’

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Type Package

Title Annotation of Metabolites from Liquid Chromatography-Mass Spectrometry Data

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Imports enviPick, ggplot2, scales, utils, stats, graphics,grDevices

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Description Automatic metabolite annotation from Liquid Chromatography-Mass Spectrometry (LC-MS and LC-MS/MS DIA) analysis.

License GPL-2

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Description

Analysis and annotation of LC-MS/MS DIA data with the use of in-house mass spectral libraries.

Usage

```
AIF(fileMS1,fileMS2,CE=0, database,rtw=7,
mzw=0.05,dmzgap=50,drt dens =20,drtgap=25,
drtsmallMS1=100,drtsmallMS2=30,dmzdensMS1=15,
dmzdensMS2=30,drtfill=5,drttotal=100,minpeakMS1=5,
minpeakMS2=3,recurs=2,weight=2,SB=3, SN=2,
minintMS1=1000,minintMS2=100,maxint=9e+09,
ion_mode="positive",ppm=TRUE,ended=6)
```

Arguments

fileMS1	A .mzXML file extension with the MS/MS experiment obtained at particular collision energy (CE).
fileMS2	A .mzXML file extension with the MS/MS experiment obtained at particular collision energy (CE).
CE	Collision energy employed for the MS/MS experiment.
database	A csv file with data arranged in columns including the names in the first row: Metabolite; Monoisotopic mass for the precursor; Mass Fragment 1; Mass Fragment 2; Mass Fragment 3... There is no need to used retention times as a constraint.
rtw	numeric. The difference between the theoretical retention time value and the experimental. Default value=3
mzw	numeric. The difference between the theoretical m/z value and the experimental. Default value=0.004
dmzgap	Arguments to be passed from enviPickwrap
dmzdensMS1	Arguments to be passed from enviPickwrap for MS1 mode
dmzdensMS2	Arguments to be passed from enviPickwrap for MS2 mode
drtgap	Arguments to be passed from enviPickwrap
drtsmallMS1	Arguments to be passed from enviPickwrap for MS1 mode
drtsmallMS2	Arguments to be passed from enviPickwrap for MS2 mode
drt dens	Arguments to be passed from enviPickwrap
drtfill	Arguments to be passed from enviPickwrap
drttotal	Arguments to be passed from enviPickwrap
minpeakMS1	Minimum number of scans that comprise a peak for MS1 mode.

minpeakMS2	Minimum number of scans that comprise a peak for MS2 mode.
recurs	Arguments to be passed from enviPickwrap
weight	Arguments to be passed from enviPickwrap
SB	Arguments to be passed from enviPickwrap
SN	Arguments to be passed from enviPickwrap
minintMS1	Arguments to be passed from enviPickwrap . Value for the MS1 mode.
minintMS2	Arguments to be passed from enviPickwrap . Value for the MS2 mode
maxint	Arguments to be passed from enviPickwrap
ion_mode	Arguments to be passed from enviPickwrap
ppm	Arguments to be passed from enviPickwrap
ended	Arguments to be passed from enviPickwrap

Value

ms1	Peak picking for MS1 level
ms2	Peak picking for MS2 level
annotation	Annotated metabolites after interrogation with library. In case a feature appears at more than one retention time, that one with highest intensity is selected.
annotationfull	Matrix with the metabolites annotated.
nodetected	List with the fragments no detected to each metabolite.
nodetectedMS2	List with the fragments no detected to each metabolite considering remaining ion molecular in MS2
RawData1	Raw scans from raw data.
RawData2	Raw scans from raw data.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)

CE.isolation("AIFpos1000-AIF.mzXML","fileposB")

#Reading the database.csv file:
# database<- read.csv("C:/database.csv")
```

```
#Processing peak-picking and annotation with default parameters  
#aif5<-AIF(fileMS,fileMS2CE5,database,CE=5, ion_mode = "positive")
```

CE.isolation

Separation of MS/MS files with regards collision energy

Description

Isolation of MS/MS events acquired at different collision energies into single files at particular collision energy

Usage

```
CE.isolation(file, output)
```

Arguments

file	A mzXML file from the LC-MS/MS experiment in positive or negative ionization mode.
output	Multiple mzXML files separated by collision energies.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)  
#Reading the file.mzXML  
#CE.isolation("AIFpos1000-AIF.mzXML", "fileposB")
```

Description

Peak picking of MS data is performed by the `enviPick` algorithm embedded. Second, it is performed a targeted extraction with a mass tolerance and m/z interval windows constraints for general peak grouping and library interrogation. Retention time might be considered as optional constraints. Library listing needs to follow the format following the example attached.

Usage

```
FullMS(file, database, rtw = 10, mzw = 0.001,  
dmzgap = 50, dmzdens = 20, drtgap = 25, drtsmall = 50,  
drt dens = 20, drtfill = 5, drttotal = 100, minpeak = 5,  
recurs = 3, weight = 1, SB = 2, SN = 1.5, minint = 1000,  
maxint = 9e+09, ion_mode = "positive",  
ppm = TRUE, ended=6)
```

Arguments

<code>file</code>	A mzXML file from the LC-MS/MS experiment in positive or negative ionization mode.
<code>database</code>	A file with data arranged in columns as follows: Molecular Formula; Retention time (optional); Neutral mass; Compound name
<code>rtw</code>	numeric. The difference between the theoretical retention time value and the experimental. Default value=3
<code>mzw</code>	numeric. The difference between the theoretical m/z value and the experimental (Da). Default value=0.004
<code>dmzgap</code>	Arguments to be passed from enviPickwrap
<code>dmzdens</code>	Arguments to be passed from enviPickwrap
<code>drtgap</code>	Arguments to be passed from enviPickwrap
<code>drtsmall</code>	Arguments to be passed from enviPickwrap
<code>drt dens</code>	Arguments to be passed from enviPickwrap
<code>drtfill</code>	Arguments to be passed from enviPickwrap
<code>drttotal</code>	Arguments to be passed from enviPickwrap
<code>minpeak</code>	Arguments to be passed from enviPickwrap
<code>recurs</code>	Arguments to be passed from enviPickwrap
<code>weight</code>	Arguments to be passed from enviPickwrap
<code>SB</code>	Arguments to be passed from enviPickwrap
<code>SN</code>	Arguments to be passed from enviPickwrap
<code>minint</code>	Arguments to be passed from enviPickwrap

maxint	Arguments to be passed from enviPickwrap
ion_mode	Arguments to be passed from enviPickwrap
ppm	Arguments to be passed from enviPickwrap
ended	Arguments to be passed from enviPickwrap

Value

ms	Annotated metabolites.
RawData1	Raw scans from raw data.
PP	Results obtained throughout enviPick algorithm performed on MS level 1.
Peaklist	Matrix with the peak picked obtained throughout enviPick algorithm.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)
#Reading the file.mzXML
# file<-fullMS.mzXML

#Reading the database.csv file:
# database<- read.csv("C:/FullMS1.csv")

#Processing peak-picking and annotation with default parameters
# FullMS_results<-FullMS(file,database, ion_mode = "positive",)

#Output:
#FullMS_results$ms
```

PeakGroup

Peak Grouping for multiple DIA files

Description

Annotated metabolites from single DIA files acquired at different collision energies are subjected to peak grouping. The function groups metabolites that are presented along the DIA files.

Usage

```
PeakGroup(aif1,aif2,aif3)
```

Arguments

aif1	Result obtained from AIF
aif2	Result obtained from AIF
aif3	Result obtained from AIF

Value

csv	A Peakgroup.csv files for each metabolite
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Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)
#Reading the file.mzXML for Full-MS scan and MS/MS.
#The files were previously separated with the SEPC.R function.

#fileMS<-"fullMS.mzXML"
#fileMS2CE5<-"fileMS2CE5.mzXML"

# Separation by collision energy
#CE.isolation("AIFpos1000-AIF.mzXML","fileposB")

#Reading the database.csv file:
# database<- read.csv("C:/database.csv")

#Processing peak-picking and annotation with default parameters

#aif5<-AIF(fileMS,fileMS2CE5,databasepos,CE=5,
#ion_mode = "positive",mzw = 0.005,rtw = 7
#,minintMS2=1,minintMS1=1)
#aif10<-AIF(fileMS,fileMS2CE10,databasepos,CE=10,
#ion_mode = "positive",rtw = 7, mzw = 0.005
#,minintMS2=1,minintMS1=1)
#aif20<-AIF(fileMS,fileMS2CE20,databasepos, CE=20,
#ion_mode = "positive",rtw = 7, mzw = 0.005,
#minintMS2=1,minintMS1=1)
```

```
# Peak grouping
#Peakgroup<-PeakGroup(aif5,aif10,aif20)
```

plot_EIC

Plot an Extracted Ion Chromatogram (EIC)

Description

Plot an Extracted Ion Chromatogram (EIC) either from processed MS1 or MS/MS file with [FullMS](#) or [AIF](#).

Usage

```
plot_EIC(fullms,peakID=333,ms=1,CE=0)
```

Arguments

fullms	A fullms or DIA file processed by FullMS or AIF .
peakID	Identity of the EIC desired to plot. The peakID is indicated in the output obtained with FullMS or AIF .
ms	numeric. MS level of EIC desired to plot.
CE	numeric. Collision energy for the file processed.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)
#Reading the file.mzXML
# file<-fullMS.mzXML

#Reading the database.csv file:
# database<- read.csv("C:/FullMS1.csv")

#Processing peak-picking and annotation with default parameters
```



```
# FullMS_results<-FullMS(file,database, ion_mode = "positive",)

#Output:
#FullMS_results$ms

#plot_EIC(fullmsposH,peakID=413)
```

ScoresDIA	<i>Statistical analysis for a pair of peak grouped metabolites from LC-MS/MS DIA analysis.</i>
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Description

Peak-to-peak Pearson correlation coefficient, peak-to-peak shape ratio and product/precursor ion intensity ratios are calculated for a product and precursor metabolites from LC-MS/MS DIA experiment.

Usage

```
ScoresDIA(input, file, ID1, ID2, CE)
```

Arguments

input	Peak grouped for a particular metabolite obtained with the PeakGroup .
file	LC-MS/MS DIA file processed by the AIF .
ID1	PeakID of the precursor ion metabolite.
ID2	PeakID of the product ion metabolite.
CE	numeric. Collision energy for the file processed.

Value

Score	Peak-to-peak Pearson correlation coefficient for a pair of EIC peaks.
IntensityRatio	Peak intensity ratio between product and precursor ion metabolite.
AssymetriRatio	Score for the chromatogram peak shape based on assymetry factor.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```

library(MetaboList)

#CE.isolation("AIFpos1000-AIF.mzXML","fileposB")

#Reading the database.csv file:
# database<- read.csv("C:/database.csv")

#Processing peak-picking and annotation with default parameters

#aif5<-AIF(fileMS,fileMS2CE5,database,CE=5, ion_mode = "positive")
#aif10<-AIF(fileMS,fileMS2CE10,database,CE=10, ion_mode = "positive")

#Peakgroup<-PeakGroup(aif5,aif10)

#Scores5<-ScoresDIA(Peakgroup$Glutamine,aif5,ID1=90, ID2 = 95,CE=5)

```

ScoresMS1

Statistical Analysis for a pair of peaks annotated for a particular metabolite.

Description

Isotopic distribution or multiple adducts of a particular compound can be evaluated by the ScoresMS1.R function that searches the output generated by the FullMS.R function for compounds annotated with multiple adducts or isotopes, calculating the isotope peak intensity ratio between monoisotopic or first isotopologue and the next one. Function calculates PPC based on intensity for both peaks at each scan forming EICs.

Usage

```
ScoresMS1(fullmspos, fullmsposb, ID1, ID2)
```

Arguments

fullmspos	Results achieved by FullMS.R function for a particular adduct.
fullmsposb	Results achieved by FullMS.R function for a particular adduct
ID1	PeakID of the peak corresponding to fileA.
ID2	PeakID of the peak corresponding to fileB.

Value

Score	Peak-to-peak Pearson correlation coefficient for a pair of EIC peaks.
IntensityRatio	Isotope peak intensity ratio between monoisotopic or first isotopologue and the next one.
AssymetriRatio	Score for the chromatogram peak shape based on assymetry factor.
name	Metabolite name.

Author(s)

Manuel D Peris Diaz

References

1. R-MetaboList: a flexible tool for metabolite extraction from high-resolution data-independent acquisition mass spectrometry analysis. *Metabolites*. Soon
2. A Survey of Orbitrap All Ion Fragmentation Analysis Assessed by an R MetaboList Package to Study Small-Molecule Metabolites. *Chromatographia*. 2018, 81, 981-994.

Examples

```
library(MetaboList)
#Reading the file.mzXML
# file<-fullMS.mzXML

#Reading the database.csv file:
# database<- read.csv("C:/FullMS1.csv")

#Processing peak-picking and annotation with default parameters
# FullMS_results<-FullMS(file,database, ion_mode = "negative",)

#ScoresMS1<-ScoresMS1(FullMS_results,FullMS_results,ID2=149,ID1=148)
```

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