Package 'flagme'

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Title Analysis of Metabolomics GC/MS Data

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Description Fragment-level analysis of gas chromatography - mass spectrometry metabolomics data

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addAMDISPeaks Add AMDIS peak detection results

Description

Reads ASCII ELU-format files (output from AMDIS) and attaches them to an already created peaksDataset object

Usage

```
addAMDISPeaks(object,fns=dir(,"[Eu][L1][Uu]"),verbose=TRUE,...)
```

Arguments

object	a peaksDataset object.
fns	character vector of same length as <code>object@rawdata</code> (user ensures the order matches)
verbose	whether to give verbose output, default TRUE
	arguments passed on to parseELU

Details

Repeated calls to parseELU to add peak detection results to the original peaksDataset object.

Value

peaksDataset object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

addChromaTOFPeaks

See Also

parseELU, peaksDataset

Examples

```
# need access to CDF (raw data) and ELU files
require(gcspikelite)
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")</pre>
```

```
# full paths to file names
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# create a peaksDataset object and add AMDIS peaks to it
pd<-peaksDataset(cdfFiles[1],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1])</pre>
```

addChromaTOFPeaks Add ChromaTOF peak detection results

Description

Reads ASCII tab-delimited format files (output from ChromaTOF) and attaches them to an already created peaksDataset object

Usage

```
addChromaTOFPeaks(object,fns=dir(,"[Tt][Xx][Tx]"),rtDivide=60,verbose=TRUE,...)
```

Arguments

object	a peaksDataset object.
fns	character vector of same length as <code>object@rawdata</code> (user ensures the order matches)
rtDivide	number giving the amount to divide the retention times by.
verbose	whether to give verbose output, default TRUE
	arguments passed on to parseChromaTOF

Details

Repeated calls to parseChromaTOF to add peak detection results to the original peaksDataset object.

Value

peaksDataset object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

parseChromaTOF, peaksDataset

Examples

```
# need access to CDF (raw data) and ChromaTOF files
require(gcspikelite)
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
# full paths to file names
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
# [not run] cTofFiles<-dir(gcmsPath,"txt",full=TRUE)
# create a peaksDataset object and add ChromaTOF peaks to it
pd<-peaksDataset(cdfFiles[1],mz=seq(50,550),rtrange=c(7.5,8.5))</pre>
```

```
# [not run] pd<-addChromTOFPeaks(pd,...)</pre>
```

betweenAlignment Data Structure for "between" alignment of many GCMS samples

Description

This function creates a "between" alignment (i.e. comparing merged peaks)

Usage

```
betweenAlignment(pD,cAList,pAList,impList,filterMin=3,gap=0.7,D=10,usePeaks=TRUE,df=30,verbose=TRUE
```

pD	a peaksDataset object
cAList	list of clusterAlignment objects, one for each experimental group
pAList	list of progressiveAlignment objects, one for each experimental group
impList	list of imputation lists
filterMin	minimum number of peaks within a merged peak to be kept in the analysis
gap	gap parameter
D	retention time penalty parameter

calcTimeDiffs

usePeaks	logical, whether to use peaks (if TRUE) or the full 2D profile alignment (if FALSE)
df	distance from diagonal to calculate similarity
verbose	logical, whether to print information

Details

betweenAlignment objects gives the data structure which stores the result of an alignment across several "pseudo" datasets. These pseudo datasets are constructed by merging the "within" alignments.

Value

betweenAlignment object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

multipleAlignment

Examples

```
require(gcspikelite)
# see multipleAlignment
```

calcTimeDiffs

Calculate retention time shifts from profile alignments

Description

This function takes the set of all pairwise profile alignments and use these to estimate retention time shifts between each pair of samples. These will then be used to normalize the retention time penalty of the signal peak alignment.

Usage

calcTimeDiffs(pd,ca.full,verbose=TRUE)

Arguments

pd	a peaksDataset object
ca.full	a clusterAlignment object, fit with
verbose	logical, whether to print out information

Details

Using the set of profile alignments,

Value

list of same length as ca.full@alignments with the matrices giving the retention time penalties.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksAlignment, clusterAlignment

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# pairwise alignment using all scans
fullca<-clusterAlignment(pd, usePeaks = FALSE, df = 100)</pre>
```

```
# calculate retention time shifts
timedf<-calcTimeDiffs(pd, fullca)</pre>
```

clusterAlignment Data Structure for a collection of all pairwise alignments of GCMS runs

Description

Store the raw data and optionally, information regarding signal peaks for a number of GCMS runs

Usage

clusterAlignment(pD,runs=1:length(pD@rawdata),timedf=NULL,usePeaks=TRUE,verbose=TRUE,...)

Arguments

рD	a peaksDataset object.
runs	vector of integers giving the samples to calculate set of pairwise alignments over.
timedf	list (length = the number of pairwise alignments) of matrices giving the expected time differences expected at each pair of peaks (used with usePeaks=TRUE, passed to peaksAlignment
usePeaks	logical, TRUE uses peakdata list, FALSE uses rawdata list for computing simi- larity.
verbose	logical, whether to print out info.
	other arguments passed to peaksAlignment

Details

clusterAlignment computes the set of pairwise alignments.

Value

clusterAlignment object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksDataset, peaksAlignment

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)
# read data, peak detection results
```

```
pdd-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
ca<-clusterAlignment(pd, gap = .5,D=.05,df=30)</pre>
```

compress

Compress an alignment object

Description

Many of the peaks are not similar. So, the set of pairwise similarity matrices can be compressed.

Usage

```
compress(object,verbose=TRUE,...)
decompress(object,verbose=TRUE,...)
```

Arguments

object	a peaksAlignment, peaksAlignment or peaksAlignment object to be com- pressed
verbose	logical, whether to print out information
	further arguments

Details

Using sparse matrix representations, a significant compression can be achieved. Here, we use the matrix.csc class of the SpareM package.

Value

an object of the same type as the input object

Author(s)

Mark Robinson

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dp

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksAlignment, clusterAlignment, progressiveAlignment

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)
# read data, peak detection results
```

```
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# pairwise alignment (it is compressed by default)
ca<-clusterAlignment(pd, usePeaks = TRUE, df = 20)
object.size(ca)</pre>
```

```
# decompress
ca<-decompress(ca)
object.size(ca)</pre>
```

Dynamic programming algorithm, given a similarity matrix

Description

This function calls C code for a bare-bones dynamic programming algorithm, finding the best cost path through a similarity matrix.

Usage

dp(M,gap=.5,big=1000000000,verbose=FALSE)

М	similarity matrix
gap	penalty for gaps
big	large value used for matrix margins
verbose	logical, whether to print out information

Details

This is a pretty standard implementation of a bare-bones dynamic programming algorithm, with a single gap parameter and allowing only simple jumps through the matrix (up, right or diagonal).

Value

list with element match with the set of pairwise matches.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

normDotProduct

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# similarity matrix
r<-normDotProduct(pd@peaksdata[[1]],pd@peaksdata[[2]])</pre>
```

```
# dynamic-programming-based matching of peaks
v<-dp(r,gap=.5)</pre>
```

eitherMatrix-class The eitherMatrix class

Description

A container to store either matrix or matrix.csc objects

Author(s)

Mark Robinson

gatherInfo

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksAlignment

gatherInfo

Gathers abundance informations from an alignment

Description

Given an alignment table (indices of matched peaks across several samples) such as that within a progressiveAlignment or multipleAlignment object, this routines goes through the raw data and collects the abundance of each fragment peak, as well as the retention times across the samples.

Usage

gatherInfo(pD, obj, newind = NULL, method = c("apex"), findmzind = TRUE, useTIC = FALSE, top = NULL, inte

Arguments

рD	a peaksDataset object, to get the abundance data from
obj	either a multipleAlignment or progressiveAlignment object
newind	list giving the
method	method used to gather abundance information, only apex implemented currently.
findmzind	logical, whether to take a subset of all m/z indices
useTIC	logical, whether to use total ion current for abundance summaries
top	only use the top top peaks
intensity.cut	percentage of the maximum intensity

Details

This procedure loops through the table of matched peaks and gathers the

Value

Returns a list (of lists) for each row in the alignment table. Each list has 3 elements:

mz	a numerical vector of the m/z fragments used
rt	a numerical vector for the exact retention time of each peak across all samples
data	matrix of fragment intensities. If useTIC=TRUE, this matrix will have a single
	row

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

imputePeaks

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# multiple alignment
ma<-multipleAlignment(pd,c(1,1),wn.gap=0.5,wn.D=.05,bw.gap=0.6,bw.D=.2,usePeaks=TRUE,filterMin=1,df=50,verbose</pre>
```

```
# gather apex intensities
d<-gatherInfo(pd,ma)</pre>
```

```
# table of retention times
nm<-list(paste("MP",1:length(d),sep=""),c("S1","S2"))
rts<-matrix(unlist(sapply(d,.subset,"rt")),byrow=TRUE,nc=2,dimnames=nm)</pre>
```

```
imputePeaks
```

Imputatin of locations of peaks that were undetected

Description

Using the information within the peaks that are matched across several runs, we can impute the location of the peaks that are undetected in a subset of runs

Usage

```
imputePeaks(pD, obj, type = 1, obj2 = NULL, filterMin = 3, verbose = TRUE)
```

imputePeaks

Arguments

рD	a peaksDataset object
obj	the alignment object, either multipleAlignment or progressiveAlignment, that is used to infer the unmatched peak locations
type	type of imputation to do, 1 for simple linear interpolation (default), 2 only works if obj2 is a clusterAlignment object
obj2	a clusterAlignment object
filterMin	minimum number of peaks within a merged peak to impute
verbose	logical, whether to print out information

Details

If you are aligning several samples and for a (small) subset of the samples in question, a peak is undetected, there is information within the alignment that can be useful in determining where the undetected peak is, based on the surrounding matched peaks. Instead of moving forward with missing values into the data matrices, this procedures goes back to the raw data and imputes the location of the apex (as well as the start and end), so that we do not need to bother with post-hoc imputation or removing data because of missing components.

We realize that imputation is prone to error and prone to attributing intensity from neighbouring peaks to the unmatched peak. We argue that this is still better than having to deal with these in statistical models after that fact. This may be an area of future improvement.

Value

list with 3 elements apex, start and end, each masked matrices giving the scan numbers of the imputed peaks.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

multipleAlignment, progressiveAlignment, peaksDataset

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:3],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:3])
# alignments
ca<-clusterAlignment(pd, gap = .5,D=.05,df=30)</pre>
```

```
pa<-progressiveAlignment(pd, ca, gap = .6, D=.1,df=30)</pre>
```

```
v<-imputePeaks(pd,pa,filterMin=1)</pre>
```

multipleAlignment-class

Data Structure for multiple alignment of many GCMS samples

Description

Store the raw data and optionally, information regarding signal peaks for a number of GCMS runs

Usage

```
multipleAlignment(pd,group,bw.gap=0.8,wn.gap=0.6,bw.D=.20,wn.D=.05,filterMin=3,lite=FALSE,usePeaks=
```

pd	a peaksDataset object
group	factor variable of experiment groups, used to guide the alignment algorithm
bw.gap	gap parameter for "between" alignments
wn.gap	gap parameter for "within" alignments
bw.D	distance penalty for "between" alignments
wn.D	distance penalty for "within" alignments
filterMin	minimum number of peaks within a merged peak to be kept in the analysis
lite	logical, whether to keep "between" alignment details (default, FALSE)
usePeaks	logical, whether to use peaks (if TRUE) or the full 2D profile alignment (if FALSE)
df	distance from diagonal to calculate similarity
verbose	logical, whether to print information
timeAdjust	logical, whether to use the full 2D profile data to estimate retention time drifts (Note: time required)
doImpute	logical, whether to impute the location of unmatched peaks

normDotProduct

Details

multipleAlignment is the data structure giving the result of an alignment across several GCMS runs.

Multiple alignments are done progressively. First, all samples with the same tg\$Group label with be aligned (denoted a "within" alignment). Second, each group will be summarized into a pseudodata set, essentially a spectrum and retention time for each matched peak of the within-alignment. Third, these "merged peaks" are aligned in the same progressive manner, here called a "between" alignment.

Value

multipleAlignment object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksDataset, betweenAlignment, progressiveAlignment

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# multiple alignment
ma<-multipleAlignment(pd,c(1,1),wn.gap=0.5,wn.D=.05,bw.gap=0.6,bw.D=.2,usePeaks=TRUE,filterMin=1,df=50,verbose</pre>
```

normDotProduct Normalized Dot Product

Description

This function calculates the similarity of all pairs of peaks from 2 samples, using the spectra similarity

Usage

normDotProduct(x1,x2,t1=NULL,t2=NULL,df=max(ncol(x1),ncol(x2)),D=100000,timedf=NULL,verbose=FALSE)

Arguments

x1	data matrix for sample 1
x2	data matrix for sample 2
t1	vector of retention times for sample 1
t2	vector of retention times for sample 2
df	distance from diagonal to calculate similarity
D	retention time penalty
timedf	matrix of time differences to normalize to. if NULL, 0 is used.
verbose	logical, whether to print out information

Details

Efficiently computes the normalized dot product between every pair of peak vectors and returns a similarity matrix. C code is called.

Value

matrix of similarities

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

dp, peaksAlignment

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

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parseChromaTOF

r<-normDotProduct(pd@peaksdata[[1]],pd@peaksdata[[2]])</pre>

parseChromaTOF Parser for ChromaTOF files

Description

Reads ASCII ChromaTOF-format files from AMDIS (Automated Mass Spectral Deconvolution and Identification System)

Usage

parseChromaTOF(fn,min.pc=.01,mz=seq(85,500),rt.cut=.008,rtrange=NULL,skip=1,rtDivide=60)

Arguments

fn	ChromaTOF filename to read.
min.pc	minimum percent of maximum intensity.
mz	vector of mass-to-charge bins of raw data table.
rt.cut	the difference in retention time, below which peaks are merged together.
rtrange	retention time range to parse peaks from, can speed up parsing if only interested in a small region (must be numeric vector of length 2)
skip	number of rows to skip at beginning of the ChromaTOF
rtDivide	multiplier to divide the retention times by (default: 60)

Details

parseChromaTOF will typically be called by addChromaTOFPeaks, not called directly.

Peaks that are detected within rt.cut are merged together. This avoids peaks which are essentially overlapping.

Fragments that are less than min.pc of the maximum intensity fragment are discarded.

Value

list with components peaks (table of spectra – rows are mass-to-charge and columns are the different detected peaks) and tab (table of features for each detection), according to what is stored in the ChromaTOF file.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

addAMDISPeaks

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
tofFiles<-dir(gcmsPath,"tof",full=TRUE)</pre>
```

parse ChromaTOF file
cTofList<-parseChromaTOF(tofFiles[1])</pre>

parseELU

Parser for ELU files

Description

Reads ASCII ELU-format files from AMDIS (Automated Mass Spectral Deconvolution and Identification System)

Usage

parseELU(f,min.pc=.01,mz=seq(50,550),rt.cut=.008,rtrange=NULL)

Arguments

f	ELU filename to read.
min.pc	minimum percent of maximum intensity.
mz	vector of mass-to-charge bins of raw data table.
rt.cut	the difference in retention time, below which peaks are merged together.
rtrange	retention time range to parse peaks from, can speed up parsing if only interested in a small region (must be numeric vector of length 2)

Details

parseELU will typically be called by addAMDISPeaks, not called directly.

Peaks that are detected within rt.cut are merged together. This avoids peaks which are essentially overlapping.

Fragments that are less than min.pc of the maximum intensity fragment are discarded.

peaksAlignment-class

Value

list with components peaks (table of spectra – rows are mass-to-charge and columns are the different detected peaks) and tab (table of features for each detection), according to what is stored in the ELU file.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

addAMDISPeaks

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# parse ELU file
eluList<-parseELU(eluFiles[1])</pre>
```

peaksAlignment-class Data Structure for pairwise alignment of 2 GCMS samples

Description

Store the raw data and optionally, information regarding signal peaks for a number of GCMS runs

Usage

```
peaksAlignment(d1,d2,t1,t2,gap=.5,D=1000,timedf=NULL,df=30,verbose=TRUE,usePeaks=TRUE,compress=TRUE
```

d1	matrix of MS intensities for 1st sample (if doing a peak alignment, this contains
	peak apexes/areas; if doing a profile alignment, this contains scan intensities.
	Rows are m/z bins, columns are peaks/scans.
d2	matrix of MS intensities for 2nd sample
t1	vector of retention times for 1st sample

t2	vector of retention times for 2nd sample
gap	gap penalty for dynamic programming algorithm
D	time penalty (on same scale as retention time differences, t1 and t2)
timedf	list (length = the number of pairwise alignments) of matrices giving the expected time differences expected at each pair of peaks (used with usePeaks=TRUE.
df	integer, how far from the diagonal to go to calculate the similarity of peaks. Smaller value should run faster, but be careful not to choose too low.
verbose	logical, whether to print out info.
usePeaks	logical, TRUE uses peakdata list, FALSE uses rawdata list for computing simi- larity.
compress	logical, whether to compress the similarity matrix into a sparse format.

Details

peaksAlignment is a hold-all data structure of the raw and peak detection data.

Value

peaksAlignment object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksDataset, clusterAlignment

Examples

see clusterAlignment, it calls peaksAlignment

peaksDataset

Description

Store the raw data and optionally, information regarding signal peaks for a number of GCMS runs

Usage

peaksDataset(fns=dir(,"[Cc][Dd][Ff]"),verbose=TRUE,mz=seq(50,550),rtDivide=60,rtrange=NULL)

Arguments

fns	character vector, filenames of raw data in CDF format.
verbose	logical, if TRUE then iteration progress information is output.
mz	vector giving bins of raw data table.
rtDivide	number giving the amount to divide the retention times by.
rtrange	retention time range to limit data to (must be numeric vector of length 2)

Details

peaksDataset is a hold-all data structure of the raw and peak detection data.

Value

peaksDataset object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

Examples

```
require(gcspikelite)
```

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
show(pd)</pre>
```

plot.peaksDataset

Description

Store the raw data and optionally, information regarding signal peaks for a number of GCMS runs

Usage

```
.plotpA(object,xlab="Peaks - run 1",ylab="Peaks - run 2",plotMatches=TRUE,matchPch=19,matchLwd=3,
matchCex=.5,matchCol="black",col=colorpanel(50,"black","blue","white"),
breaks=seq(0,1,length=51),...)
```

```
.plotcA(object,alignment=1,...)
```

object	a peaksDataset, peaksAlignment or clusterAlignment object.	
runs	for peaksDataset only: set of run indices to plot	
mzind	for peaksDataset only: set of mass-to-charge indices to sum over (default, all)	
mind	for peaksDataset only: matrix of aligned indices	
plotSampleLabel	ls	
	for peaksDataset only: logical, whether to display sample labels	
calcGlobalMax	for peaksDataset only: logical, whether to calculate an overall maximum for scaling	
peakCex	character expansion factor for peak labels	
plotPeaks	for peaksDataset only: logical, whether to plot hashes for each peak	
plotPeakBoundaries		
	for peaksDataset only: logical, whether to display peak boundaries	
plotPeakLabels	for peaksDataset only: logical, whether to display peak labels	
plotMergedPeakLabels		
	for peaksDataset only: logical, whether to display 'merged' peak labels	
mlwd	for peaksDataset only: line width of lines indicating the alignment	
usePeaks	for peaksDataset only: logical, whether to plot alignment of peaks (otherwise, scans)	
plotAcrossRuns	for peaksDataset only: logical, whether to plot across peaks when unmatched peak is given	

overlap	for peaksDataset only: logical, whether to plot TIC/XICs overlapping
rtrange	for peaksDataset only: vector of length 2 giving start and end of the X-axis
cols	for peaksDataset only: vector of colours (same length as the length of runs)
thin	for peaksDataset only: when usePeaks=FALSE, plot the alignment lines every thin values
max.near	for peaksDataset only: where to look for maximum
how.near	for peaksDataset only: how far away from max.near to look
scale.up	for peaksDataset only: a constant factor to scale the TICs
plotMatches	for peaksDataset only: logical, whether to plot matches
xlab	for peaksAlignment and clusterAlignment only: x-axis label
ylab	for peaksAlignment and clusterAlignment only: y-axis label
matchPch	for peaksAlignment and clusterAlignment only: match plotting character
matchLwd	for peaksAlignment and clusterAlignment only: match line width
matchCex	for peaksAlignment and clusterAlignment only: match character expansion factor
matchCol	for peaksAlignment and clusterAlignment only: match colour
col	$for {\tt peaksAlignment} and {\tt clusterAlignment} only: vector of colours for colourscale$
breaks	for $peaksAlignment$ and $clusterAlignment$ only: vector of breaks for colourscale
alignment	for peaksAlignment and clusterAlignment only: the set of alignments to plot
	further arguments passed to the plot or image command

Details

For peakDataset objects, each TIC is scale to the maximum value (as specified by the how.near and max.near values). The many parameters gives considerable flexibility of how the TICs can be visualized.

For peakAlignment objects, the similarity matrix is plotted and optionally, the set of matching peaks. clusterAlignment objects are just a collection of all pairwise peakAlignment objects.

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

plotImage, peaksDataset

Examples

```
require(gcspikelite)
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)
# read data
pd<-peaksDataset(cdfFiles[1:3],mz=seq(50,550),rtrange=c(7.5,8.5))
# image plot
plot(pd,rtrange=c(7.5,8.5),plotPeaks=TRUE,plotPeakLabels=TRUE)</pre>
```

plotImage

Plot of images of GCMS data

Description

Image plots (i.e. 2D heatmaps) of raw GCMS profile data

Usage

plotImage(object,run=1,rtrange=c(11,13),main=NULL,mzrange=c(50,200),SCALE=log2,...)

Arguments

object	a peaksDataset object
run	index of the run to plot an image for
rtrange	vector of length 2 giving start and end of the X-axis (retention time)
main	main title (auto-constructed if not specified)
mzrange	vector of length 2 giving start and end of the Y-axis (mass-to-charge ratio)
SCALE	function called to scale the data (default: log2)
	further arguments passed to the image command

Details

For peakDataset objects, each TIC is scale to the maximum value (as specified by the how.near and max.near values). The many parameters gives considerable flexibility of how the TICs can be visualized.

For peakAlignment objects, the similarity matrix is plotted and optionally, the set of matching peaks. clusterAlignment objects are just a collection of all pairwise peakAlignment objects.

Author(s)

Mark Robinson

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References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

plot, peaksDataset

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data
pd<-peaksDataset(cdfFiles[1],mz=seq(50,550),rtrange=c(7.5,8.5))</pre>
```

```
# image plot
plotImage(pd,run=1,rtrange=c(7.5,8.5),main="")
```

progressiveAlignment-class

Data Structure for progressive alignment of many GCMS samples

Description

Performs a progressive peak alignment (clustalw style) of multiple GCMS peak lists

Usage

progressiveAlignment(pD,cA,D=1000,gap=.5,verbose=TRUE,usePeaks=TRUE,df=30,compress=TRUE)

рD	a peaksDataset object
cA	a clusterAlignment object
D	retention time penalty
gap	gap parameter
verbose	logical, whether to print information
usePeaks	logical, whether to use peaks (if TRUE) or the full 2D profile alignment (if FALSE)
df	distance from diagonal to calculate similarity
compress	logical, whether to store the similarity matrices in sparse form

Details

The progressive peak alignment we implemented here for multiple GCMS peak lists is analogous to how clustalw takes a set of pairwise sequence alignments and progressively builds a multiple alignment. More details can be found in the reference below.

Value

progressiveAlignment object

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksDataset, multipleAlignment

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
ca<-clusterAlignment(pd, gap = .5,D=.05,df=30)
pa<-progressiveAlignment(pd, ca, gap = .6, D=.1,df=30)</pre>
```

rmaFitUnit

Fits a robust linear model (RLM) for one metabolite

Description

Using rlm from MASS, this procedure fits a linear model using all the fragments

Usage

rmaFitUnit(u,maxit=5,mzEffect=TRUE,cls=NULL,fitSample=TRUE,fitOrCoef=c("coef","fit"),TRANSFORM=log

rmaFitUnit

Arguments

u	a metabolite unit (list object with vectors mz and rt for m/z and retention times, respectively and a data element giving the fragmentxsample intensitity matrix)
maxit	maximum number of iterations (default: 5)
mzEffect	logical, whether to fit m/z effect (default: TRUE)
cls	class variable
fitSample	whether to fit individual samples (alternative is fit by group)
fitOrCoef	whether to return a vector of coefficients (default: "coef"), or an rlm object ("fit")
TRANSFORM	function to transform the raw data to before fitting (default: log2)

Details

Fits a robust linear model.

Value

list giving elements of fragment and sample coefficients (if fitOrCoef="coef") or a list of elements from the fitting process (if fitOrCoef="fit")

Author(s)

Mark Robinson

References

Mark D Robinson (2008). Methods for the analysis of gas chromatography - mass spectrometry data *PhD dissertation* University of Melbourne.

See Also

peaksAlignment, clusterAlignment

Examples

require(gcspikelite)

```
# paths and files
gcmsPath<-paste(find.package("gcspikelite"),"data",sep="/")
cdfFiles<-dir(gcmsPath,"CDF",full=TRUE)
eluFiles<-dir(gcmsPath,"ELU",full=TRUE)</pre>
```

```
# read data, peak detection results
pd<-peaksDataset(cdfFiles[1:2],mz=seq(50,550),rtrange=c(7.5,8.5))
pd<-addAMDISPeaks(pd,eluFiles[1:2])</pre>
```

```
# pairwise alignment using all scans
fullca<-clusterAlignment(pd, usePeaks = FALSE, df = 100)</pre>
```

rmaFitUnit

calculate retention time shifts
timedf<-calcTimeDiffs(pd, fullca)</pre>

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