Package 'metabCombiner'

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Title Method for Combining LC-MS Metabolomics Feature Measurements

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Description This package aligns LC-HRMS metabolomics datasets acquired from biologically similar specimens analyzed under similar, but not necessarily identical, conditions. Peak-picked and simply aligned metabolomics feature tables (consisting of m/z, rt, and per-sample abundance measurements, plus optional identifiers & adduct annotations) are accepted as input. The package outputs a combined table of feature pair alignments, organized into groups of similar m/z, and ranked by a similarity score. Input tables are assumed to be acquired using similar (but not necessarily identical) analytical methods.

Depends R (>= 4.0)

Imports dplyr (>= 1.0), methods, mgcv, caret, S4Vectors, stats, utils, rlang, graphics, matrixStats, tidyr

Suggests knitr, rmarkdown, testthat, BiocStyle

BugReports https://www.github.com/hhabra/metabCombiner/issues

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Collate 'adjustData.R' 'batchCombine.R' 'calcScores.R' 'check_pars.R' 'classes.R' 'combinerCheck.R' 'compare_strings.R' 'data.R' 'detectFields.R' 'evaluateParams.R' 'fit_model.R' 'form.R' 'generics.R' 'labelRows.R' 'metabCombine.R' 'metabCombiner_package_doc.R' 'metabData.R' 'methods-featdata.R' 'methods-metabCombiner.R' 'methods-metabData.R' 'mzGroup.R' 'params.R' 'plot_fit.R' 'resolveRows.R' 'selectAnchors.R' 'updateTables.R' 'write2file.R' 'zzz.R'

VignetteBuilder knitr

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Description

This retrieves user-assigned adduct annotations from one or all constituent datasets of a metabCombiner object

Usage

```
adductData(object, data = NULL)
## S4 method for signature 'metabCombiner'
adductData(object, data = NULL)
```

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Arguments

object metabCombiner object

data dataset identifier to extract information from; if NULL, extracts information

from all datasets

Value

data frame of adduct annotations

Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")

p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")

##retrieve all adduct data
adducts <- adductData(p.comb, data = NULL)

##retrieve adduct data from p30
adducts <- adductData(p.comb, data = "p30")</pre>
```

adjustData

Process and Filter Metabolomics Feature Lists

Description

adjustData contains a set of pre-analysis steps for processing LC-MS metabolomics feature tables individually

Usage

```
adjustData(Data, misspc, measure, rtmin, rtmax, zero, duplicate)
```

Arguments

Data a metabData object.

misspc Numeric. Threshold missingness percentage for analysis.

measure Character. Choice of central abundance measure; either "median" or "mean".

rtmin Numeric. Minimum retention time for analysis.

rtmax Numeric. Maximum retention time for analysis.

zero Logical. Whether to consider zero values as missing.

duplicate Ordered numeric pair (m/z, rt) tolerance parameters for duplicate feature search.

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Details

The pre-analysis adjustment steps include: 1) Restriction to a feature retention time range $rtmin \le rt \le rtmax 2$) Removal of features with missingness percentage exceeding misspc 3) Removal of duplicate metabolomics features.

After processing, abundance quantile (Q) values are calculated between 0 & 1 for the remaining features, as ranked by the measure argument, unless provided by the user.

Value

Updated metabData object. The data field is processed by the listed steps and stats list updated to contain feature statistics.

See Also

```
metabData, filterRT
```

batchCombine

Stepwise Multi-batch LC-MS Alignment

Description

This is a method for aligning multiple batches of a single metabolomics experiment in a stepwise manner using the metabCombiner workflow. The input is a list of metabData objects corresponding to the batch data frames arranged in sequential order (i.e. batch 1,2,...,N), and parameter lists for each step; the output is an aligned feature table and a metabCombiner object composed from the input batches.

Usage

```
batchCombine(
  batches,
  binGap = 0.005,
  fitMethod = "gam",
  means = list(mz = TRUE, rt = TRUE, Q = TRUE),
  union = FALSE,
  anchorParam = selectAnchorsParam(),
  fitParam = fitgamParam(),
  scoreParam = calcScoresParam(B = 30),
  reduceParam = reduceTableParam()
)
```

Arguments

batches list of metabData objects corresponding to each LC-MS batch

binGap numeric parameter used for grouping features by m/z. See ?mzGroup for more

details.

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fitMethod RT spline-fitting method, either "gam" or "loess"

means logical. Option to take average m/z, rt, and/or Q from metabCombiner. May be

a 3-length vector, single value (TRUE/FALSE), or a list with names "mz", "rt",

"Q" as names.

union logical. If FALSE, only feature present in all batches will be in the final result.

If TRUE, features missing in at least one batch are included. The mean m/z, RT,

and Q values imputed for matching in each step.

anchorParam list of parameter values for selectAnchors() function fitParam list of parameter values for fit_gam() or fit_loess()

scoreParam list of parameter values for calcScores()
reduceParam list of parameter values for reduceTable()

Details

Retention time drifting is commonly observed in large-scale LC-MS experiments in which samples are analyzed in multiple batches. Conventional LC-MS pre-processing approaches may effectively align features detected in samples from within a single batch, but fail in many cases to account for inter-batch drifting, leading to misaligned features.

batchCombine assumes that each batch has been previously processed separately using conventional LC-MS preprocessing approaches (e.g. XCMS), and can be represented as a data frame. Each batch data feature table must be filtered and formatted as a metabData object and the batches must be arranged as a list in sequential order of acquisition.

batchCombine applies the metabCombine wrapper function to successive pairs of metabolomics batches in a stepwise manner. Each iteration consists of the key steps in the package workflow (feature m/z grouping, anchor selection, retention time spline fitting, pairwise scoring, & table reduction). The first two batches are aligned together, then the combined results are aligned with the third batch, and so forth. Parameters for each sub-method are arranged in list format, with their respective defaults (e.g. fitgamParam() lists the default values for the fit_gam function).

Following each iteration, m/z, rt, and Q values from the combined dataset may be averaged to use for comparison with the next batch's feature quantitative descriptors, if the means argument is set to TRUE; if set to FALSE, feature information is drawn from the latter of the previously combined batches, identical to the manner in which id & adduct descriptors are drawn.

Value

object metabCombiner object of the final alignment; x is set to the penultimate batch

and y is set to the final batch

table combined feature table consisting of feature descriptor values followed by per-

sample abundances and extra columns

Note

batchCombine is designed for aligning multi-batch datasets, i.e. where each batch is acquired in a roughly identical manner. It is not for disparately acquired LC-MS datasets (e.g. from different instruments, chromatographic systems, laboratories, etc.).

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See Also

metabCombine

Examples

```
#identically formatted batches in list form
data(metabBatches)
#obtain list of metabData objects
batchdata <- lapply(metabBatches, metabData, samples = "POOL",</pre>
                     extra = "SAMP", zero = TRUE)
#recommended: give each batch dataset a unique name
names(batchdata) <- paste("exb", seq_along(batchdata), sep = "")</pre>
#customize main workflow parameter lists
saparam <- selectAnchorsParam(tolmz = 0.002, tolQ = 0.2, tolrtq = 0.1)</pre>
fgparam <- fitgamParam(k = 20, iterFilter = 1)</pre>
csparam \leftarrow calcScoresParam(A = 70, B = 35, C = 0.3)
rtparam <- reduceTableParam(minScore = 0.5, maxRTerr = 0.33)</pre>
#run batchCombine program
combinedRes <- batchCombine(batches = batchdata, binGap = 0.0075,</pre>
               means = list('mz' = TRUE, 'rt' = FALSE, 'Q' = FALSE),
               anchorParam = saparam, fitParam = fgparam,
               scoreParam = csparam, reduceParam = rtparam)
#aligned table results & metabCombiner object results
cTable <- combinedRes$table
object <- combinedRes$object</pre>
#if names were set earlier, the names should be returned by this
datasets(object)
```

calcScores

Compute Feature Similarity Scores

Description

Calculates a pairwise similarity (between 0 & 1) between all grouped features in metabCombiner object. The similarity score calculation is described in scorePairs.

Usage

```
calcScores(
  object,
  A = 75,
```

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```
B = 10,
C = 0.25,
groups = NULL,
fit = c("gam", "loess"),
mzshift = FALSE,
mzfit = mzfitParam(),
useAdduct = FALSE,
adduct = 1.25,
usePPM = FALSE,
brackets_ignore = c("(", "[", "{"}")
```

Arguments

object	metabCombiner object.
A	Numeric weight for penalizing m/z differences.
В	Numeric weight for penalizing differences between fitted & observed retention times
С	Numeric weight for differences in Q (abundance quantiles).
groups	integer. Vector of feature groups to score. If set to NULL (default), will compute scores for all feature groups.
fit	Character. Choice of fitted rt model, "gam" or "loess."
mzshift	Logical. If TRUE, shifts the m/z values (mzx) before scoring.
mzfit	List of parameters for shifting m/z values; see ?mzfitParam
useAdduct	logical. Option to penalize mismatches in (non-empty, non-bracketed) adduct column annotations.
adduct	numeric. If useAdduct is TRUE, divides score of mismatched, non-empty and non-bracked adduct column labels by this value.
usePPM	logical. Option to use relative (as opposed to absolute) m/z differences in score computations.
brackets_ignor	e

Details

This function updates the rtProj, score, rankX, and rankY columns in the combinedTable report. First, using the RT mapping model computed in the previous steps, rtx values are projected onto rty. Then similarity scores are calculated based on m/z, rt (rtProj vs rty), and Q differences, with multiplicative weight penalties A, B, and C.

ignored according to this argument

If useAdduct = TRUE, bracketed adduct character strings of these types will be

If the datasets contain representative set of shared identities (idx = idy), evaluateParams provides some guidance on appropriate A, B, and C values to use. In testing, the best values for A should lie between 50 and 120, according to mass accuracy; if using ppm (usePPM = TRUE), the suggested range is between 0.01 and 0.05. B should be between 5 and 15 depending on fitting accuracy (higher if datasets processed under roughly identical conditions); C should vary between 0 and 1, depending on sample similarity. See examples below.

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Some input datasets exhibit systematic m/z shifts

If using adduct information (useAdduct = TRUE), the score is divided by the numeric adduct argument if non-empty and non-bracketed adduct values do not match. Be sure that adduct annotations are accurate before using this functionality.

Value

metabCombiner object with updated combinedTable. rtProj column will contain fitted retention times determined from previously computed model; score will contain computed pairwise similarity scores of feature pairs; rankX & rankY are the integer ranks of scores for x & y features in descending order.

See Also

```
evaluateParams, scorePairs
```

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)</pre>
p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 1, family = "gaussian")</pre>
#example: moderate m/z deviation, accurate rt fit, high sample similarity
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.8, useAdduct = FALSE,
         groups = NULL, fit = "gam", usePPM = FALSE)
cTable <- combinedTable(p.comb) #to view results
#example 2: high m/z deviation, moderate rt fit, low sample similarity
p.comb \leftarrow calcScores(p.comb, A = 50, B = 8, C = 0.2)
#example 3: low m/z deviation, poor rt fit, moderate sample similarity
p.comb \leftarrow calcScores(p.comb, A = 120, B = 5, C = 0.5)
#example 4: using ppm for mass deviation; note different A value
p.comb <- calcScores(p.comb, A = 0.05, B = 14, C = 0.5, usePPM = TRUE)
#example 5: limiting to specific m/z groups 1-1000
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.5, groups = seq(1,1000))
#example 6: using adduct information
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.5, useAdduct = TRUE,
                     adduct = 1.25)
```

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calcScoresParam

List calcScores Defaults

Description

List of default parameters for score calculation step of main package workflow. See help(calcScores) or ?calcScores for details.

Usage

```
calcScoresParam(
    A = 75,
    B = 10,
    C = 0.25,
    fit = "gam",
    groups = NULL,
    usePPM = FALSE,
    useAdduct = FALSE,
    adduct = 1.25,
    brackets_ignore = c("(", "[", "{"}")")
```

Arguments

Α	m/z difference specific weight; default: 75
В	RT prediction error specific weight; default: 10
С	Q difference specific weight; default: 0.25
fit	choice of fitted model ("gam" or "loess"); default: "gam"
groups	choice of m/z groups to score
usePPM	choice to use PPM for m/z differences; default: FALSE
useAdduct	choice to use adduct strings in scoring; default: FALSE
adduct	value divisor for mismatched adduct strings; default: 1.25
brackets_ignore	
	bracket types for ignoring string comparisons

Value

list of calcScores parameters

See Also

```
calcScores, metabCombine
```

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Examples

```
cs_param <- calcScoresParam(A = 60, B = 15, C = 0.3)
cs_param <- calcScoresParam(A = 0.1, B = 20, C = 0.2, usePPM = TRUE)</pre>
```

combineData

Obtain All Feature Data

Description

Obtain all meta-data (m/z, RT, Q, id, adduct) alongside their respective sample (+ extra) values for aligned features. This is a (quasi)merge of the /code/linkcombinedTable and /code/linkfeatData tables and methods.

Usage

```
combineData(object)
## S4 method for signature 'metabCombiner'
combineData(object)
```

Arguments

object

metabCombiner object

Value

A data.frame containing meta-data columns as well as sample + extra columns for each of the constituent data sets.

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")

p.comb <- metabCombiner(p30, p20)
p.comb.table <- combineData(p.comb)</pre>
```

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Obtain Feature Alignment Report

Description

Obtain constructed table reporting every feature pair alignment.

Usage

```
combinedTable(object)

## S4 method for signature 'metabCombiner'
combinedTable(object)
```

Arguments

idx

object metabCombiner object.

Value

Feature Pair Alignment report data frame. The columns of the report are as follows:

Identities of features from dataset X

idy	Identities of features from dataset Y
mzx	m/z values of features from dataset X
mzy	m/z values of features from dataset Y
rtx	retention time values of features from dataset X
rty	retention time values of features from dataset Y
rtProj	model-projected (X->Y) retention times values
Qx	abundance quantile values of features from dataset X
Qy	abundance quantile values of features from dataset Y
group	m/z feature group of feature pairing

computed similarity scores of feature pairing
rankX ranking of pairing score for X dataset features
rankY ranking of pairing score for Y dataset features

adductX adduct label of features from dataset X adductY adduct label of features from dataset Y

... Sample and extra columns from both datasets X & Y

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Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")

p.comb <- metabCombiner(p30, p20)
p.comb.table <- combinedTable(p.comb)</pre>
```

combinerCheck

Obtain Errors for metabCombiner Object Checks

Description

This function stores and returns a customized error message when checking the validity of certain objects.

Usage

```
combinerCheck(errNo, type, error = "stop")
```

Arguments

errNo integer error code.

type character object type (either "combinedTable", "metabCombiner" or "metab-

Data")

error character. If "stop", gives an error message; if "warning", provides a warning

message; if "silent", returns silently

Details

In certain functions, an object must be checked for correctness. A metabData must have a properly formatted dataset with the correct column names & types.A metabCombiner must have properly formatted combinedTable, with expected names and columns. If one of these conditions is not met, a non-zero numeric code is returned and this function is used to print a specific error message corresponding to the appropriate object and error code.

Value

A customized error message for specific object check.

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crossValFit

Cross Validation for Model Fits

Description

Helper function for fit_gam() & fit_loess(). Determines optimal value of k basis functions for Generalized Additive Model fits or span for loess fits from among user-defined choices, using a 10-fold cross validation minimizing mean squared error.

Usage

```
crossValFit(
  rts,
  fit,
  vals,
  bs,
  family,
  m,
  method,
  optimizer,
  control,
  message,
  ...
)
```

Arguments

rts	data.frame of ordered pair retention times
fit	Either "gam" for GAM fits, or "loess" for loess fits
vals	numeric vector: k values for GAM fits, spans for loess fits. Best value chosen by 10-fold cross validation.
bs	character. Choice of spline method, either "bs" or "ps"
family	character. Choice of mgcv family; see: ?mgcv::family.mgcv
m	integer. Basis and penalty order for GAM; see ?mgcv::s
method	character. Smoothing parameter estimation method; see: ?mgcv::gam
optimizer	character. Method to optimize smoothing parameter; see: ?mgcv::gam
control	control parameters for loess fits; see: ?loess.control
message	Option to print message indicating function progress
	Other arguments passed to mgcv::gam.

Value

Optimal parameter value as determined by 10-fold cross validation

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datasets

Obtain Dataset IDs

Description

Each dataset in a metabCombiner object is represented by a character identifier. The datasets slot contains all these ids in a single vector, which can be obtained in sequential order with this accessor method

Usage

```
datasets(object, list = FALSE)
## S4 method for signature 'metabCombiner'
datasets(object, list = FALSE)
```

Arguments

object metabCombiner object

list logical, option to return in list format (TRUE) vs character vector format (FALSE)

Value

character vector of dataset identifiers

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")

p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")

##datasets extraction: expect "p30", "p20"
sets <- datasets(p.comb, list = FALSE)</pre>
```

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detectFields Detect metabData Input Columns

Description

This function ensures that metabolomics datasets used as inputs for the program possess all of the required fields, plus any optional columns that may appear in the final report table.

Usage

```
detectFields(Data, table, mz, rt, id, adduct, samples, extra, Q)
```

Arguments

Data	a metabData object.
table	data frame containing metabolomics features or path to metabolomics data file.
mz	Character name(s) / regular expressions associated with data column containing m/z values. The first column whose name contains this expression will be selected for analysis.
rt	Character name(s) / regular expression associated with data column containing retention time values. The first column whose name contains this expression will be selected for analysis.
id	Character name(s) or regular expression associated with data column containing metabolomics feature identifiers. The first column whose name contains this expression will be selected for analysis.
adduct	Character name(s) or regular expression associated with data column containing adduct, formula, or additional annotations. The first column whose name contains this expression will be selected for analysis.
samples	Character names of columns containing sample values. All numeric columns containing these keywords are selected for analysis. If no keywords given, searches for longest stretch of numeric columns remaining.
extra	Character names of columns containing additional feature information, e.g. non-analyzed sample values. All columns containing these keywords are selected for analysis.
Q	Character name(s) or regular expression associated with numeric feature abundance quantiles.

Value

an initialized and formatted metabData object.

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evaluateParams	Evaluate Similarity Score Parameters
----------------	--------------------------------------

Description

This function provides a method for guiding selection of suitable values for A, B, & C weight arguments in the calcScores method, based on the similarity scores of shared identified compounds. Datasets must have at least one identity in common (i.e. idx = idy, case-insensitive), and preferably more than 10.

Usage

```
evaluateParams(
  object,
  A = seq(60, 150, by = 10),
  B = seq(6, 15),
  C = seq(0.1, 0.5, by = 0.1),
  fit = c("gam", "loess"),
  usePPM = FALSE,
  minScore = 0.5,
  penalty = 5,
  groups = NULL,
  brackets_ignore = c("(", "[", "{"}")")
```

Arguments

object	metabCombiner object	
Α	Numeric weights for penalizing m/z differences.	
В	Numeric weights for penalizing differences between fitted & observed retention times	
С	Numeric weight for differences in Q (abundance quantiles).	
fit	Character. Choice of fitted rt model, "gam" or "loess."	
usePPM	logical. Option to use relative parts per million (ppm) as opposed to absolute) m/z differences in score computations.	
minScore	numeric minimum score to count towards objective function calculation for known matching features (idx = idy) and mismatches.	
penalty	numeric. Subtractive mismatch penalty.	
groups	integer. Vector of feature groups to score. If set to NULL (default), will compute scores for all feature groups.	
brackets_ignore		
	bracketed identity and adduct character strings of these types will be ignored	

according to this argument

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Details

This uses an objective function, based on the accurate and inaccurate alignments of shared preidentified compounds. For more details, see: objective.

Value

A data frame with the following columns:

A m/z weight values

B rt weight values

C Q weight values

totalScore objective function evaluation of (A,B,C) weights

Note

In contrast to calcScores function, A, B, & C take numeric vectors as input, as opposed to constants. The total number of rows in the output will be equal to the products of the lengths of these input vectors

See Also

```
calcScores, objective
```

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)

p.comb <- selectAnchors(p.comb, windx = 0.03, windy = 0.02)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 2)

#example 1
scores <- evaluateParams(p.comb, A = seq(60,100,10), B = seq(10,15), C = 0.5, minScore = 0.7, penalty = 10)

##example 2: limiting to groups 1-2000
scores <- evaluateParams(p.comb, minScore = 0.5, groups = seq(1,2000))</pre>
```

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featData

Obtain Feature Metadata

Description

This method retrieves all feature meta-data or that of one data set. The rowIDs identically correspond to the rows from the combinedTable data frame.

Usage

```
featData(object, data = NULL)
## S4 method for signature 'metabCombiner'
featData(object, data = NULL)
```

Arguments

object a metabCombiner object data character dataset identifier

Details

metabCombiner objects organized metabolomics feature information in the "featData" slot. This table and method is primarily useful for alignment analyses involving three or more data sets.

Value

data frame of feature metadata from one or all datasets

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")

p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")

#full metadata extraction
fdata <- featData(p.comb, data = NULL)

#single dataset feature information extraction
fdata <- featData(p.comb, data = "p20")</pre>
```

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filterAnchors

Filter Outlier Ordered Pairs

Description

Helper function for fit_gam & fit_loess. It filters the set of ordered pairs using the residuals calculated from multiple GAM / loess fits.

Usage

```
filterAnchors(
  rts,
 fit,
 vals,
 outlier,
 coef,
  iterFilter,
 prop,
  bs,
 m,
  family,
 method,
 optimizer,
  control,
 message,
)
```

Arguments

rts	Data frame of ordered retention time pairs.
fit	Either "gam" for GAM fits, or "loess" for loess fits
vals	numeric values: k values for GAM fits, spans for loess fits
outlier	Thresholding method for outlier dection. If "MAD", the threshold is the mean absolute deviation (MAD) times coef; if "boxplot", the threshold is coef times IQR plus 3rd quartile of a model's absolute residual values.
coef	numeric (> 1) multiplier for determining thresholds for outliers (see outlier argument)
iterFilter	integer number of outlier filtering iterations
prop	numeric. A point is excluded if deemed a residual in more than this proportion of fits. Must be between $0 \& 1$.
bs	character. Choice of spline method from mgcv; either "bs" or "ps"
m	integer. Basis and penalty order for GAM; see ?mgcv::s
family	character. Choice of mgcv family; see: ?mgcv::family.mgcv

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method	character. Smoothing parameter estimation method; see: ?mgcv::gam
optimizer	character. Method to optimize smoothing parameter; see: ?mgcv::gam
control	control parameters for loess fits; see: ?loess.control
message	Option to print message indicating function progress
	other arguments passed to mgcv::gam.

Value

anchor rts data frame with updated weights.

filtered	Retrieve Filtered Features	

Description

Returns a data frame of metabolomics features eliminated in the metabData step. Features are returned based on the specific filter used for their elimination (RT, missingness, or duplicates).

Usage

```
filtered(object, type = c("rt", "duplicates", "missing"))
## S4 method for signature 'metabData'
filtered(object, type = c("rt", "missing", "duplicates"))
```

Arguments

object metabData object
type one of three filter types used for feature removal

Value

data frame of features removed due to specified filter

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filterRT	Filter Features by Retention Time Range	

Description

Restricts input metabolomics feature table in metabData object to a range of retention times defined by rtmin & rtmax.

Usage

```
filterRT(data, rtmin, rtmax)
```

Arguments

data	formatted metabolomics data frame.
rtmin	lower range of retention times for analysis. If " \min ", defaults to minimum observed retention time
rtmax	upper range of retention times for analysis. If "max", defaults to maximum

observed retention time.

Details

Retention time restriction is often recommended to aid the analysis of comparable metabolomics datasets. The beginning and end of a chromatogram typically contain features that do not correspond with true biological compounds derived from the sample. rtmin and rtmax should be set slightly before and slightly after the first and last commonly observed metabolites, respectively.

Value

A data frame of metabolomics features, limited to time window $rtmin \le rt \le rtmax$)

Description

List of default parameters for GAM fitting step of main package workflow, which can be used as input for the wrapper functions. See help(fit_gam) or ?fit_gam for more details.

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Usage

```
fitgamParam(
    useID = FALSE,
    k = seq(10, 20, 2),
    iterFilter = 2,
    outlier = "MAD",
    coef = 2,
    prop = 0.5,
    weights = 1,
    bs = "bs",
    family = "scat",
    m = c(3, 2),
    method = "REML",
    rtx = c("min", "max"),
    rty = c("min", "max"),
    optimizer = "newton",
    message = TRUE
)
```

Arguments

useID	choice of preserving identity-based anchors; default: FALSE
k	values for GAM basis dimension k
iterFilter	number of outlier filtering iterations; default: 2
outlier	outlier filtering method (either "MAD" (mean absolute deviation) or "boxplot"); default: "MAD" $$
coef	outlier filtering coefficient; default: 2
prop	minimum proportion of fits in which a point can be a flagged outlier; default: 0.5
weights	optional supplied weights to individual points; default: 1
bs	choice of spline type ("bs" or "ps"); default: "bs"
family	choice of family ("scat" or "gaussian"); default: "scat"
m	basis and penalty order; default: $c(3,2)$
method	smoothing parameter estimation method; default: "REML"
rtx	ordered pair of endpoints for rtx; default: ("min", "max")
rty	ordered pair of endpoints for rty; default: ("min", "max")
optimizer	numerical optimization for GAM; default: "newton"
message	option to print progress message; default: TRUE

Value

list of fit_gam parameters

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See Also

```
fit_gam, metabCombine
```

Examples

fitloessParam

List fitLoess Defaults

Description

List of default parameters for loess fitting step of main package workflow, See help(fit_loess) or ?fit_loess for more details.

Usage

```
fitloessParam(
  useID = FALSE,
  spans = seq(0.2, 0.3, by = 0.02),
  outlier = "MAD",
  coef = 2,
  iterFilter = 2,
  prop = 0.5,
  weights = 1,
  message = TRUE,
  rtx = c("min", "max"),
  rty = c("min", "max"),
  control = loess.control(surface = "direct", iterations = 10)
)
```

Arguments

```
choice of preserving identity-based anchors; default: FALSE
useID
                   values for span parameter which controls degree of smoothing
spans
outlier
                   outlier filtering method (either "MAD" or "boxplot"); default: "MAD"
                   outlier filtering coefficient; default: 2
coef
iterFilter
                   number of outlier filtering iterations; default: 2
                   minimum proportion of fits where a point can be a flagged outlier; default: 0.5
prop
weights
                   optional supplied weights to individual points; default: 1
message
                   option to print progress message; default: TRUE
rtx
                   ordered pair of endpoints for rtx; default: ("min", "max")
                   ordered pair of endpoints for rty; default: ("min", "max")
rty
control
                   loess-specific control parameters; see: ?loess.control
```

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Value

list of fit_loess parameters:

See Also

```
fit_loess, metabCombine
```

Examples

```
fitParam <- fitloessParam(spans = c(0.2,0.25,0.3), outlier = "boxplot", iterFilter = 3, coef = 1.5, message = FALSE, control = loess.control(iterations = 4))
```

 ${\tt fit_gam}$

Fit RT Projection Model With GAMs

Description

Fits a (penalized) basis splines curve through a set of ordered pair retention times, modeling one set of retention times (rty) as a function on the other set (rtx). Outlier filtering iterations are performed first, then with the remaining points, the best value of parameter k is selected through 10-fold cross validation.

Usage

```
fit_gam(
 object,
 useID = FALSE,
 k = seq(10, 20, 2),
  iterFilter = 2,
 outlier = c("MAD", "boxplot"),
  coef = 2,
 prop = 0.5,
 weights = 1,
 bs = c("bs", "ps"),
 m = c(3, 2),
  family = c("scat", "gaussian"),
 method = "REML",
  rtx = c("min", "max"),
  rty = c("min", "max"),
 optimizer = "newton",
 message = TRUE,
)
```

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Arguments

object	a metabCombiner object.
useID	logical. If set to TRUE, matched ID anchors detected from previous step will never be flagged as outliers.
k	integer k values controlling the dimension of the basis of the GAM fit (see: ?mgcv::s). Best value chosen by 10-fold cross validation.
iterFilter	integer number of outlier filtering iterations to perform
outlier	Thresholding method for outlier dection. If "MAD", the threshold is the mean absolute deviation (MAD) times coef; if "boxplot", the threshold is coef times IQR plus 3rd quartile of a model's absolute residual values.
coef	numeric (> 1) multiplier for determining thresholds for outliers (see outlier argument)
prop	numeric. A point is excluded if deemed a residual in more than this proportion of fits. Must be between $0\ \&\ 1.$
weights	Optional user supplied weights for each ordered pair. Must be of length equal to number of anchors (n) or a divisor of $(n+2)$.
bs	character. Choice of spline method from mgcv, either "bs" (basis splines) or "ps" (penalized basis splines)
m	integer. Basis and penalty order for GAM; see ?mgcv::s
family	character. Choice of mgcv family; see: ?mgcv::family.mgcv
method	character smoothing parameter estimation method; see: ?mgcv::gam
rtx	ordered pair of endpoints for rtx; if "max" or "min", gives the maximum or minimum rtx, respectively, as model endpoints for rtx $\frac{1}{2}$
rty	ordered pair of endpoints for rty; if "max" or "min", gives the maximum or minimum rtx, respectively, as model endpoints for rty
optimizer	character. Method to optimize smoothing parameter; see: ?mgcv::gam
message	Option to print message indicating function progress
	Other arguments passed to mgcv::gam.

Details

A set of ordered pair retention times must be previously computed using selectAnchors(). The minimum and maximum retention times from both input datasets are included in the set as ordered pairs (min_rtx, min_rty) & (max_rtx, max_rty). The weights argument initially determines the contribution of each point to the model fits; they are equally weighed by default, but can be changed using an n+2 length vector, where n is the number of ordered pairs and the first and last of the weights determines the contribution of the min and max ordered pairs; by default, all weights are initially set to 1 for equal contribution of each point.

The model complexity is determined by k. Multiple values of k are allowed, with the best value chosen by 10 fold cross validation. Before this happens, certain ordered pairs are removed based on the model errors. In each iteration, a GAM is fit using each selected value of k. Depending on the outlier argument, a point is "removed" from the model (i.e. its corresponding weight set to 0) if its residual is above the threshold for a proportion of fitted models, as determined by prop. If an

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anchor is an "identity" (idx = idy, detected in the selectAnchors by setting useID to TRUE), then setting useID here prevents its removal.

Other arguments, e.g. family, m, optimizer, bs, and method are GAM specific parameters from the mgcv R package. The family option is currently limited to the "scat" (scaled t) and "gaussian" families; scat family model fits are more robust to outliers than gaussian fits, but compute much slower. Type of splines are currently limited to basis splines ("bs" or "ps").

Value

metabCombiner with a fitted GAM model object

See Also

```
selectAnchors,fit_loess,
```

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb = metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)
p.comb = selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
anchors = getAnchors(p.comb)
#version 1: using faster, but less robust, gaussian family
p.comb = fit_gam(p.comb, k = c(10, 12, 15, 17, 20), prop = 0.5,
    family = "gaussian", outlier = "MAD", coef = 2)
#version 2: using slower, but more robust, scat family
p.comb = fit_gam(p.comb, k = seq(12,20,2), family = "scat",
                     iterFilter = 1, coef = 3, method = "GCV.Cp")
#version 3 (with identities)
p.comb = selectAnchors(p.comb, useID = TRUE)
anchors = getAnchors(p.comb)
p.comb = fit_gam(p.comb, useID = TRUE, k = seq(12,20,2), iterFilter = 1)
#version 4 (using identities and weights)
weights = ifelse(anchors$labels == "I", 2, 1)
p.comb = fit_gam(p.comb, useID = TRUE, k = seq(12,20,2),
                     iterFilter = 1, weights = weights)
#version 5 (using boxplot-based outlier detection
p.comb = fit_gam(p.comb, k = seq(12,20,2), outlier = "boxplot", coef = 1.5)
#to preview result of fit_gam
plot(p.comb, pch = 19, outlier = "h", xlab = "CHEAR Plasma (30 min)",
    ylab = "Red-Cross Plasma (20 min)", main = "Example GAM Fit")
```

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fit_loess

Fit RT Projection Model With LOESS

Description

Fits a local regression smoothing spline through a set of ordered pair retention times. modeling one set of retention times (rty) as a function on the other set (rtx). Filtering iterations of high residual points are performed first. Multiple acceptable values of span can be used, with one value selected through 10-fold cross validation.

Usage

```
fit_loess(
  object,
  useID = FALSE,
  spans = seq(0.2, 0.3, by = 0.02),
  outlier = c("MAD", "boxplot"),
  coef = 2,
  iterFilter = 2,
  prop = 0.5,
  weights = 1,
  rtx = c("min", "max"),
  rty = c("min", "max"),
  message = TRUE,
  control = loess.control(surface = "direct", iterations = 10)
)
```

Arguments

object	a metabCombiner object.
useID	logical. If set to TRUE, matched ID anchors detected from previous step will never be flagged outliers.
spans	numeric span values (between 0 & 1) used for loess fits
outlier	Thresholding method for outlier dection. If "MAD", the threshold is the mean absolute deviation (MAD) times coef; if "boxplot", the threshold is coef times IQR plus 3rd quartile of a model's absolute residual values.
coef	numeric (> 1) multiplier for determining thresholds for outliers (see outlier argument)
iterFilter	integer number of outlier filtering iterations to perform
prop	numeric. A point is excluded if deemed a residual in more than this proportion of fits. Must be between 0 & 1.

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weights	Optional user supplied weights for each ordered pair. Must be of length equal to number of anchors (n) or a divisor of $(n+2)$
rtx	ordered pair of endpoints for rtx; if "max" or "min", gives the maximum or minimum rtx, respectively, as model endpoints for rtx
rty	ordered pair of endpoints for rty; if "max" or "min", gives the maximum or minimum rtx, respectively, as model endpoints for rty
message	Option to print message indicating function progress
control	control parameters for loess fits; see: ?loess.control

Value

metabCombiner object with model slot updated to contain a fitted loess model

See Also

```
selectAnchors,fit_gam
```

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb = metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)
p.comb = selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
#version 1
p.comb = fit_loess(p.comb, spans = seq(0.2,0.3,0.02), iterFilter = 1)
#version 2 (using weights)
anchors = getAnchors(p.comb)
weights = c(2, rep(1, nrow(anchors)), 2) #weight = 2 to boundary points
p.comb = fit_loess(p.comb, spans = seq(0.2,0.3,0.02), weights = weights)
#version 3 (using identities)
p.comb = selectAnchors(p.comb, useID = TRUE, tolmz = 0.003)
p.comb = fit_loess(p.comb, spans = seq(0.2,0.3,0.02), useID = TRUE)
#to preview result of fit_loess
plot(p.comb, fit = "loess", xlab = "CHEAR Plasma (30 min)",
     ylab = "Red-Cross Plasma (20 min)", pch = 19,
    main = "Example fit_loess Result Fit")
```

30 getAnchors

getAnchors

Get Ordered Retention Time Pairs

Description

Returns the data frame of feature alignments used to anchor the retention time projection model, constructed by selectAnchors.

Usage

```
getAnchors(object)
## S4 method for signature 'metabCombiner'
getAnchors(object)
```

Arguments

object

metabCombiner object

Value

Data frame of anchor features

See Also

selectAnchors

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red")

p.comb <- metabCombiner(p30, p20)
p.comb <- selectAnchors(p.comb, windx = 0.05, windy = 0.03)
anchors <- getAnchors(p.comb)</pre>
```

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getCoefficients

Obtain Last-Used Score Coefficients

Description

Provides the last used weight arguments from calcScores() function. Returns empty list if calcScores() has not yet been called.

Usage

```
getCoefficients(object)
## S4 method for signature 'metabCombiner'
getCoefficients(object)
```

Arguments

object

metabCombiner object

Value

A list of the last used weight parameters:

A Specific weight penalizing feature m/z differences

B Specific weight penalizing retention time projection error

C Specific weight penalizing differences in abundance quantiles

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red")

p.comb <- metabCombiner(p30, p20)
p.comb <- selectAnchors(p.comb, windx = 0.05, windy = 0.04, tolrtq = 0.15)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 1, family = "gaussian")
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.5)

getCoefficients(p.comb)</pre>
```

32 getExtra

getData

Get Processed Dataset

Description

The metabData constructor creates a formatted dataset from the input, which may be accessed using this method.

Usage

```
getData(object)
## S4 method for signature 'metabData'
getData(object)
```

Arguments

object

metabData object

Value

Single Metabolomics Data Frame

Examples

```
data(plasma30)
p30 <- metabData(plasma30, samples = "CHEAR")
data <- getData(p30)</pre>
```

getExtra

Get Extra Data Column Names

Description

Get Extra Data Column Names

Usage

```
getExtra(object, data = NULL)
## S4 method for signature 'metabCombiner'
getExtra(object, data = NULL)
## S4 method for signature 'metabData'
getExtra(object)
```

getModel 33

Arguments

object metabCombiner or metabData object

data dataset identifier for metabCombiner objects

Value

character vector of extra column names

Examples

```
data(plasma30)
p30 <- metabData(plasma30, samples = "CHEAR", extra = "Red")
getExtra(p30)</pre>
```

getModel

Get Fitted RT Model

Description

Returns the last fitted RT projection model from a metabCombiner object of type "gam" or "loess".

Usage

```
getModel(object, fit = c("gam", "loess"))
## S4 method for signature 'metabCombiner'
getModel(object, fit = c("gam", "loess"))
```

Arguments

object metabCombiner object

fit Choice of model, "gam" or "loess"

Value

nonlinear retention time fit object

See Also

```
fit_gam, fit_loess
```

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Examples

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.005)
p.comb <- selectAnchors(p.comb, tolrtq = 0.15, tolQ = 0.2, windy = 0.02)
p.comb <- fit_gam(p.comb, iterFilter = 1, k = 20, family = "gaussian")
p.comb <- fit_loess(p.comb, iterFilter = 1, spans = 0.2)
model.gam <- getModel(p.comb, fit = "gam")
model.loess <- getModel(p.comb, fit = "loess")</pre>
```

getSamples

Get Sample Names From metabCombiner or metabData Object

Description

Returns the sample names from one of the two datasets used in metabCombiner analysis, denoted as 'x' or 'y.'

Usage

```
getSamples(object, data = NULL)
## S4 method for signature 'metabCombiner'
getSamples(object, data = NULL)
## S4 method for signature 'metabData'
getSamples(object)
```

Arguments

object metabCombiner or metabData object
data dataset identifier for metabCombiner objects

Value

character vector of sample names. For metabCombiner objects these may come from the 'x' dataset (if data = "x") or the 'y' dataset (if data = "y").

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)</pre>
```

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```
p.comb <- metabCombiner(xdata = p30, ydata = p20)

getSamples(p30)
getSamples(p.comb, data = "x") #equivalent to previous
getSamples(p20)
getSamples(p.comb, data = "y") #equivalent to previous</pre>
```

getStats

Get Object Statistics

Description

Prints out a list of object-specific statistics for both metabCombiner and metabData objects

Usage

```
getStats(object)
## S4 method for signature 'metabCombiner'
getStats(object)
## S4 method for signature 'metabData'
getStats(object)
```

Arguments

object

metabCombiner or metabData object

Value

list of object-specific statistics

Methods (by class)

• metabCombiner: Method for 'metabCombiner' object

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)

getStats(p30) #metabData stats

p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.005)
p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)</pre>
```

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```
p.comb <- fit_gam(p.comb, iterFilter = 1, k = 20)
getStats(p.comb) #metabCombiner stats</pre>
```

idData

Retrieve Feature Identities

Description

This retrieves user-assigned feature identities from one or all constituent datasets of a metabCombiner object

Usage

```
idData(object, data = NULL)
## S4 method for signature 'metabCombiner'
idData(object, data = NULL)
```

Arguments

object metabCombiner object

dataset identifier to extract information from; if NULL, extracts information

from all datasets

Value

data frame of feature identities

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")
p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")

##retrieve all ids
ids <- idData(p.comb, data = NULL)

##retrieve ids from p30
ids <- idData(p.comb, data = "p30")</pre>
```

identityAnchorSelection

Select Matching Ids as Anchors

Description

This is an optional helper function for selectAnchors. Uses identities to guide selection of ordered retention time pairs. If useID option is set to TRUE, it will select pairs of features with matching ID character strings before proceeding with iterative anchor selection.

Usage

identityAnchorSelection(cTable, windx, windy, useID, brackets)

Arguments

cTable	data frame, contains only feature ids, mzs, rts, Qs, & labels
windx	numeric positive retention time exclusion window in X dataset
windy	numeric positive retention time exclusion window in Y dataset
useID	logical. Operation proceeds if TRUE, terminates otherwise.
brackets	If useID = TRUE, bracketed identity strings of the types included in this argument will be ignored

Details

Identity anchors are allowed to violate constraints of m/z, Q, and rtq difference tolerances, and will not be removed if they fall within a rt exclusion window of other features. If a name appears more than once, only the pair with the highest relative abundance is selected.

Value

combinedTable with updated anchor labels

See Also

selectAnchors

38 isMetabCombiner

isCombinedTable

Determine combinedTable Validity

Description

Checks whether input object is a valid metabData.Returns an integer code if invalid. Function is used alongside combinerCheck.

Usage

isCombinedTable(object)

Arguments

object

Any R object.

Details

a proper combinedTable must have these characteristics to be deemed valid for metabCombiner operations:

- 1) It must be a data.frame with at least 16 columns and at least 1 row
- & "adducty" in this exact order

 2) The first 16 columns must be of closer "numeric" "character" "character" "numeric" "numeri

2) The first 16 columns must be named "rowID", "idx", "idy", "mzx", "mzy", "rtx", "rty", "rtProj", "Qx", "Qy", "group", "score", "r

- 3) The first 16 columns must be of class: "numeric" "character", "character", "numeric", "numeric", "numeric", "numeric", "integer", "integer", "integer", "integer", "integer", "character", "character", "character", "character", "integer", "i
- 4) The group column must have no missing or negative values

Failing any one of these criteria causes an error

Value

0 if object is a valid Combiner Table; an integer code otherwise

isMetabCombiner

Determine if object is a valid metabCombiner object

Description

Checks whether input object is a valid metabCombiner.Returns an integer code if invalid. Function is used alongside combinerCheck.

Usage

isMetabCombiner(object)

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Arguments

object

Any R object.

Value

0 if object is a valid metabData object; an integer code otherwise

isMetabData

Determine validity of input metabData object

Description

Checks whether input object is a valid metabData.Returns an integer code if invalid. Function is used alongside combinerCheck.

Usage

isMetabData(object)

Arguments

object

Any R object

Value

0 if object is a valid metabData object; an integer code otherwise.

iterativeAnchorSelection

Iterative Selection of Ordered Pairs

Description

This is a helper function for selectAnchors. Anchors are iteratively selected from highly abundant feature pairs, subject to feature m/z, rt, & Q constraints set by the user.

Usage

```
iterativeAnchorSelection(cTable, windx, windy, swap = FALSE)
```

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Arguments

cTable	data frame, contains only feature ids, mzs, rts, Qs, & labels
windx	numeric positive retention time exclusion window in X dataset.
windy	numeric positive retention time exclusion windown in Y dataset.
swap	logical. When FALSE, searches for abundant features in dataset X, complemented by dataset Y features; when TRUE, searches for abundant features in dataset Y, complemented by dataset X features.

Value

data frame of anchor feature alignments.

See Also

selectAnchors

labelRows

Annotate and Remove Report Rows

Description

This is a method for annotating removable, conflicting, and identity-matched feature pair alignment (FPA) rows in the combinedTable report. Simple thresholds for score, rank, retention time error and delta score can computationally reduce the set of possible FPAs to the most likely feature matches. FPAs falling within some small delta score or mz/rt of the top-ranked pair are organized into subgroups to facilitate inspection. Automated reduction to 1-1 pairs is also possible with this function.

reduceTable behaves identically to labelRows, but with a focus on automated table reduction. Rank threshold defaults in reduceTable are also stricter than in labelRows.

Usage

```
labelRows(
  object,
  useID = FALSE,
  minScore = 0.5,
  maxRankX = 3,
  maxRankY = 3,
  delta = 0.1,
  method = c("score", "mzrt"),
  maxRTerr = 10,
  resolveConflicts = FALSE,
  rtOrder = TRUE,
  remove = FALSE,
  balanced = TRUE,
  brackets_ignore = c("(", "[", "{"}")
```

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```
reduceTable(
  object,
  useID = FALSE,
  maxRankX = 2,
  maxRankY = 2,
  minScore = 0.5,
  delta = 0.1,
  method = c("score", "mzrt"),
  maxRTerr = 10,
  rtOrder = TRUE,
  brackets_ignore = c("(", "[", "{"}"))
```

Arguments

object Either a metabCombiner object or combinedTable

useID option to annotate identity-matched strings as "IDENTITY"

minScore numeric minimum allowable score (between 0 & 1) for metabolomics feature

pair alignments

maxRankX integer maximum allowable rank for X dataset features.

maxRankY integer maximum allowable rank for Y dataset features.

delta numeric score or mz/rt distances used to define subgroups. If method = "score",

a value (between 0 & 1) score difference between a pair of conflicting FPAs. If method = "mzrt", a length 4 numeric: (m/z, rt, m/z, rt) tolerances, the first pair

for X dataset features and the second pair for Y dataset features.

method Conflict detection method. If equal to "score" (default), assigns a conflict sub-

group if score of lower-ranking FPA is within some tolerance of higher-ranking FPA. If set to "mzrt", assigns a conflicting subgroup if within a small m/z & rt

distance of the top-ranked FPA.

maxRTerr numeric maximum allowable error between model-projected retention time (rt-

Proj) and observed retention time (rty)

resolveConflicts

logical option to computationally resolve conflicting rows to a final set of 1-1

feature pair alignments

rtOrder logical. If resolveConflicts set to TRUE, then this imposes retention order con-

sistency on rows deemed "RESOLVED" within subgroups.

remove Logical. Option to keep or discard rows deemed removable.

balanced Logical. Optional processing of "balanced" groups, defined as groups with an

equal number of features from input datasets where all features have a 1-1 match.

brackets_ignore

character. If useID = TRUE, bracketed identity strings of the types in this argu-

ment will be ignored

42 labelRows

Details

metabCombiner initially reports all possible feature pairings in the rows of the combinedTable report. Most of these are misalignments that require removal. This function is used to automate this reduction process by labeling rows as removable or conflicting, based on certain conditions, and is performed after computing similarity scores.

A label may take on one of four values:

a) "": No determination made b) "IDENTITY": an alignment with matching identity "idx & idy" strings c) "REMOVE": a row determined to be a misalignment d) "CONFLICT": competing alignments for one or multiple shared features

The labeling rules are as follows:

1) Groups determined to be 'balanced': label rows with rankX > 1 & rankY > 1 "REMOVE" irrespective of delta criteria 2) Rows with a score < minScore: label "REMOVE" 3) Rows with rankX > maxRankX and/or rankY > maxRankY: label "REMOVE" 4) Conflicting subgroup assignment as determined by method & delta arguments. Conflicting alignments following outside delta thresholds: labeled "REMOVE". Otherwise, they are assigned a "CONFLICT" label and subgroup number. 5) If useID argument set to TRUE, rows with matching idx & idy strings are labeled "IDENTITY". These rows are not changed to "REMOVE" or "CONFLICT" irrespective of subsequent criteria.

Value

updated combinedTable or metabCombiner object. The table will have three new columns:

labels characterization of feature alignments as described subgroup conflicting subgroup number of feature alignments alt alternate subgroup for rows in multiple feature pair conflicts

Examples

```
#required steps prior to function use
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)</pre>
p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 1)</pre>
p.comb \leftarrow calcScores(p.comb, A = 90, B = 14, C = 0.5)
##applies labels, but maintains all rows
p.comb <- labelRows(p.comb, maxRankX = 2, maxRankY = 2, maxRTerr = 0.5,</pre>
                     delta = 0.1, resolveConflicts = FALSE, remove = FALSE)
##automatically resolve conflicts and filter to 1-1 feature pairs
p.comb.2 <- labelRows(p.comb, resolveConflicts = FALSE, remove = FALSE)</pre>
#this is identical to the previous command
p.comb.2 <- reduceTable(p.comb)</pre>
```

labelRowsParam 43

labelRowsParam

List labelRows & reduceTable Defaults

Description

List of default parameters for combinedTable row annotation and removal. See help(labelRows) or ?labelRows for more details. reduceTableParam loads parameters for the more automated reduceTable function

Usage

```
labelRowsParam(
  useID = FALSE,
 maxRankX = 3,
 maxRankY = 3,
 minScore = 0.5,
 method = "score",
 delta = 0.1,
 maxRTerr = 10,
  resolveConflicts = FALSE,
  rtOrder = TRUE,
  remove = FALSE,
 balanced = TRUE,
  brackets_ignore = c("(", "[", "{")
)
reduceTableParam(
  useID = FALSE,
 maxRankX = 2,
 maxRankY = 2,
 minScore = 0.5,
 maxRTerr = 10,
 delta = 0.1,
  rtOrder = TRUE,
 method = "score",
 brackets_ignore = c("(", "[", "{")
)
```

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Arguments

useID option to annotate identity-matched strings as IDENTITY; default: FALSE

maxRankX maximum rank allowable for X features
maxRankY maximum rank allowable for Y features
minScore minimum score threshold; default: 0.5

method thresholding method for subgroup detection ("score" or "mzrt"); default: "score" delta score distance or mz/rt difference tolerances for subgrouping; default: 0.1

maxRTerr maximum allowable difference between predicted RT (rtProj) & observed RT

(rty); default: 10 minutes

resolveConflicts

logical. If TRUE, automatically resolves subgroups to 1-1 feature pair align-

ments

rtOrder logical. If TRUE and resolveConflicts is TRUE, imposes retention order condi-

tion on paired alignments

remove option to eliminate rows determined as removable; default: FALSE

balanced option to reduce balanced groups; default: TRUE

brackets_ignore

bracket types for ignoring string comparisons

Value

list of labelRows parameters

See Also

labelRows, metabCombine, batchCombine, reduceTable

Examples

metabBatches

Three LC-MS Metabolomics Batch Datasets

Description

An example multi-batch LC-MS metabolomics analysis of human plasma, used to demonstrate batchCombine. Due to the large size of the full experimental data, only three of the batches are loaded here with a subset of the samples and features from each batch.

Usage

data(metabBatches)

metabCombine 45

Format

A list containing three identically formatted data frames

metabCombine

metabCombiner Wrapper Function

Description

metabCombine wraps the five main metabCombiner workflow steps into a single wrapper function. Parameter list arguments organize program parameters by constituent package functions.

Usage

```
metabCombine(
  xdata,
 ydata,
 binGap = 0.005,
  xid = NULL,
 yid = NULL,
 means = list(mz = FALSE, rt = FALSE, Q = FALSE),
  fitMethod = "gam",
  rtOrder = TRUE,
  union = FALSE,
  impute = FALSE,
  anchorParam = selectAnchorsParam(),
  fitParam = fitgamParam(),
  scoreParam = calcScoresParam(),
  labelParam = labelRowsParam()
)
```

Arguments

xdata	metabData object. One of two datasets to be combined.
ydata	metabData object. One of two datasets to be combined.
binGap	numeric parameter used for grouping features by m/z . See ? mz Group for more details.
xid	character identifier of xdata. If xdata is a metabData, assigns a new ID for this dataset; if xdata is a metabCombiner, must be assigned to one of the existing dataset IDs. See details for more information.
yid	character identifier of ydata. If ydata is a metabData, assigns a new ID for this dataset; if ydata is a metabCombiner, must be assigned to one of the existing dataset IDs. See details for more information.
means	logical. Option to take average m/z, rt, and/or Q from metabComber. May be a vector (length = 3), single value (TRUE/FALSE), or a list with names "mz", "rt", "Q" as names.

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fitMethod	RT spline-fitting method, either "gam" or "loess"
rtOrder	logical. If set to TRUE, retention order consistency expected when resolving conflicting alignments for metabCombiner object inputs.
union	logical. Option to include non-matched features in final combinedTable results
impute	logical. If TRUE, imputes the mean mz/rt/Q values for missing features in metabCombiner object inputs before use in alignment (not recommended for disparate data alignment); if FALSE, features with missing information are dropped.
anchorParam	list of parameter values for selectAnchors() function
fitParam	list of parameter values for fit_gam() or fit_loess()
scoreParam	list of parameter values for calcScores()
labelParam	list of parameter values for labelRows()

Details

The five main steps in metabCombine are 1) m/z grouping & combined table construction, 2) selection of ordered pair RT anchors, 3) nonlinear spline (Basis Spline GAM or LOESS) fitting to predict RTs, 4) score calculation and feature pair alignment ranking, 5) combined table row annotation and reduction. metabData arguments xdata & ydata and m/z grouping binGap are required for step 1.

Steps 2-5 are handled by anchors, fit, scores, & labels, respectively, with lists containing the argument values for each step expected for these arguments. selectAnchorsParam, fitgamParam, fitloessParam, calcScoresParam, & labelRowsParam load the default program values of selectAnchors, fit_gam, fit_loess, calcScores & labelRows, respectively. These program arguments should be modified as necessary for the datasets used for analysis.

By default, the RT fitting method (fitMethod) is set to "gam", which means the argument fit is a list of parameters for fit_gam; if the (fitMethod) argument is set to "loess", then the fit argument expects a list of fit_loess parameters.

Value

a metabCombiner object following complete analysis

See Also

selectAnchorsParam, fitgamParam, calcScoresParam, labelRowsParam, fitloessParam

Examples

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metabCombiner

Form a metabCombiner object.

Description

This constructs an object of type metabCombiner from a pair of metabolomics datasets, formatted as either metabData (single-dataset class) or metabCombiner (combined-dataset class). An initial table of possible feature pair alignments is constructed by grouping features into m/z groups controlled by the binGap argument

Usage

```
metabCombiner(
  xdata,
  ydata,
  binGap = 0.005,
  xid = NULL,
  yid = NULL,
  means = list(mz = FALSE, rt = FALSE, Q = FALSE),
  rtOrder = TRUE,
  impute = FALSE
)
```

Arguments

xdata	metabData or metabCombiner object
ydata	metabData or metabCombiner object
binGap	numeric parameter used for grouping features by m/z. See ?mzGroup for more details.
xid	character. If xdata is a metabData, assigns a new identifier for this dataset; if xdata is a metabCombiner, selects one of the existing dataset IDs to represent xdata. See details for more information.

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yid character. If ydata is a metabData, assigns a new identifier for this dataset; if

ydata is a metabCombiner, selects one of the existing dataset IDs to represent

ydata. See details for more information.

means logical. Option to take average m/z, rt, and/or Q from metabComber. May be a

vector (length = 3), a single value (TRUE/FALSE), or a list with names "mz",

"rt", "Q" as names.

rt0rder logical. If set to TRUE, retention order consistency expected when resolving

conflicting alignments for metabCombiner object inputs.

impute logical. If TRUE, imputes the mean mz/rt/Q values for missing features in

metabCombiner object inputs before use in alignment (not recommended for disparate data alignment); if FALSE, features with missing information are dropped.

Details

This function serves as a constructor of the metabCombiner combined dataset class and the entry point to the main workflow for pairwise dataset alignment. Two arguments must be specified, xdata and ydata, which must be either metabData or metabCombiner objects. There are four scenarios listed here:

- 1) If xdata & ydata are metabData objects, a new metabCombiner object is constructed with an alignment of this pair. New character identifiers are assigned to each dataset (xid & yid, respectively); if these are unassigned, then "1" and "2" will be their respective ids. xdata & ydata will be the active "dataset x" and "dataset y" used for the paired alignment.
- 2) If xdata is a metabCombiner and ydata is a metabData, then the result is the existing metabCombiner xdata augmented by an additional dataset, ydata. One set of meta-data (id, m/z, rt, Q, adduct labels) from xdata is used for alignment with the respective information from ydata, which is controlled by the xid argument; see the datasets method for extracting existing dataset ids. A new identifier yid is assigned to ydata, which must be distinct from the current dataset identifier.
- 3) If xdata is a metabData and ydata is a metabCombiner, then a similar process to #2 occurs, with xdata augmented to the existing ydata object and one of the constitutent dataset's meta-data is accessed, as controlled by the yid argument. One major difference is that rts of ydata serve as the "reference" or dependent variable in the spline-fitting step.
- 4) If xdata and ydata are both metabCombiner objects, the resulting metabCombiner object aligns information from both combined datasets. As before, one set of values contained in xdata (specified by xid argument) is used to align to the values from ydata (controlled by yid argument). The samples and extra columns are concatenated from all datasets.

For metabCombiner object inputs, the full workflow (selectAnchors, fit_gam/fit_loess, calcScores, labelRows) must be performed before further alignment. If not completed already, features are pared down to 1-1 alignments via the resolveConflicts approach (see: help(resolveRows)). Features may not be used more than twice and will be removed if they are detected as duplicates.

The mean of the numeric fields (m/z, rt, Q) from all constituent datasets can be used in alignment in place of values from a single dataset. These are controlled by the means argument. By default this is a list value with "mz", "rt" and "Q" as names, but may also accept a single logical or a length-3 logical vector. If set to a single logical value, then all three fields are averaged (TRUE) or not averaged (FALSE). If a three-length argument is supplied (e.g. c(TRUE, FALSE), then the values correspond to m/z, rt, and Q respectively. RT averaging is generally not recommended for disparate data alignment.

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If missing features have been incorporated into the metabCombiner, they an be imputed using the average m/z, rt, and Q values for that feature in datasets in which it is present by setting impute to TRUE. Likewise, this option is not recommended for disparate data alignment.

Value

a metabCombiner object constructed from xdata and ydata, with features grouped by m/z according to the binGap argument.

Note

If using a metabCombiner object as input, only one row is allowed per feature corresponding to its first appearance. It is strongly recommended to reduce the table to 1-1 paired matches prior to aligning it with a new dataset.

Examples

metabCombiner-class

'metabCombiner' Combined Metabolomics Dataset Class

Description

This is the main object for the metabCombiner package workflow. This object holds a combined feature table, along with a retention time warping model, the ordered pair anchors used to generate this model, important information organized by dataset, and key object statistics.

Slots

combinedTable data frame displaying all feature pair alignments, combining measurements of all possible shared compounds

```
featData data frame of feature metadata (id, m/z, rt, Q, adduct) anchors data frame of feature pairs used for RT warping model model list containing the last fitted nonlinear model(s) datasets list of constituent datasets from xdata & ydata inputs xy current X & Y datasets

nonmatched list of data frames consisting of nonmatched features
```

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coefficients list of last used A,B,C similarity weight values samples list of sample name vectors from input datasets extra list of extra column name vectors from input datasets stats set of useful metabCombiner statistics

metabData

Constructor for the metabData object.

Description

This is a constructor for objects of type metabData.

Usage

```
metabData(
  table,
 mz = "mz"
  rt = "rt"
  id = "id",
  adduct = "adduct",
  samples = NULL,
  Q = NULL,
 extra = NULL,
  rtmin = "min",
  rtmax = "max",
 misspc = 50,
 measure = c("median", "mean"),
 zero = FALSE,
  duplicate = opts.duplicate()
)
```

Arguments

table	Path to file containing feature table or data.frame object containing features
mz	Character name(s) or regular expression associated with data column containing m/z values. The first column whose name contains this expression will be selected for analysis.
rt	Character name(s) or regular expression associated with data column containing retention time values. The first column whose name contains this expression will be selected for analysis.
id	Character name(s) or regular expression associated with data column containing metabolomics feature identifiers. The first column whose name contains this expression will be selected for analysis.
adduct	Character name(s) or regular expression associated with data column containing adduct or chemical formula annotations. The first column whose name contains this expression will be selected for analysis.

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samples	Character name(s) or regular expression associated with data columns. All numeric columns whose names contain these keywords are selected for analysis. If no keywords given, program searches longest stretch of remaining numeric columns.
Q	Character name(s) or regular expression associated with numeric feature abundance quantiles. If NULL, abundance quantiles are calculated from sample intensities.
extra	Character names of columns containing additional feature information, e.g. non-analyzed sample values. All columns containing these keywords selected and will be displayed in the final output.
rtmin	Numeric. Minimum retention time for analysis.
rtmax	Numeric. Maximum retention time for analysis.
misspc	Numeric. Threshold missingness percentage for analysis.
measure	Central sample abundance measure, either "median" or "mean".
zero	Logical. Whether to consider zero values as missing.
duplicate	list of duplicate feature removal parameters. (see: opts.duplicate)

Details

Processed metabolomics feature table must contain columns for m/z, rt, and numeric sample intensities. Some optional fields such as identity id and adduct label columns may also be supplied. Non-analyzed columns can be included into the final output by specifying the names of these columns in the extra argument. All required arguments are checked for validity (e.g. no negative m/z or rt values, each column is used at most once, column types are valid, etc...).

Following this is a pre-analysis filtering of rows that are either: 1) Outside of a specified retention time range (rtmin,rtmax), 2) Missing in excess of misspc percent of analyzed samples, or 3) deemed duplicates by small pairwise <m/z, rt> differences. See: opts.duplicate on duplicate feature removal

Remaining features are ranked by abundance quantiles, Q, using a central measure, either "median" or "mean." Alternatively, the abundance quantiles column can be specified in the argument Q.

Value

An object of class metabData containing the specific information specified by mz,rt, samples, id, adduct, Q, and extra arguments, and adjusted by pre-processing steps.

Examples

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```
#equivalent to above
p30 <- metabData(plasma30, id = "id", samples = "CHEAR", extra = "Red")

#analyzing Red Cross samples with retention time limitations (0.5-17.5min)
p20 <- metabData(plasma20, samples = "Red", rtmin = 0.5, rtmax = 17.5)
data = getData(p20)
range(data$rt)

#using regular expressions for field searches
p30 <- metabData(plasma30, id = "identity|id|ID", samples = ".[3-5]$")
getSamples(p30)  #should print all column names ending in .3, .4, .5</pre>
```

metabData-class

'metabData' Single Metabolomics Dataset Class

Description

This class is designed to process and format input metabolomics feature tables. It stores the information from individual metabolomics datasets, including the formatted feature table, sample names, and feature statistics.

Slots

data formatted metabolomics data frame.
samples character vector of analyzed sample names
extra character vector of non-analyzed columns names
stats A list of dataset statistics
filtered A list of filtered dataset features

mzData

Retrieve m/z Values

Description

This retrieves feature m/z values from one or all constituent datasets of a metabCombiner object. Alternatively, the average m/z value can be retrieved.

Usage

```
mzData(object, data = NULL, value = c("obs", "mean"))
## S4 method for signature 'metabCombiner'
mzData(object, data = NULL, value = c("observed", "mean"))
```

mzFit 53

Arguments

object metabCombiner object

dataset identifier to extract information from; if NULL, extracts data frame in-

formation from all datasets

value Either "obs" (observed - default option) or "mean" value

Value

```
data frame of m/z values (if NULL) or single vector of m/z values
data(plasma30) data(plasma20)
p30 <- metabData(head(plasma30,500), samples = "CHEAR") p20 <- metabData(head(plasma20,500),
samples = "Red") p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")
##retrieve all m/z mzd <- mzData(p.comb, data = NULL)
##retrieve m/z from p30 mzd <- mzData(p.comb, data = "p30")
##retrieve mean m/z mzd <- mzData(p.comb, value = "mean")
```

mzFit

Compute m/z Shift Model

Description

Generates a GAM model for correcting systematic m/z shifts observed between a pair of LC-MS data sets.

Usage

```
mzFit(object, fit = mzfitParam(), plot = TRUE, ...)
```

Arguments

object metabCombiner object

fit List of m/z shift parameter values; see ?mzfitParam

plot Logical. Option to plot the m/z shift model.

... other arguments to be passed to plot

Details

Correcting for systematic m/z shifts improves the scores for feature pair alignments, yielding more accurate match hypotheses. This function generates a basis spline curve, modeling the m/z shift (mzy - mzx) as a function of m/z (mzx). Selected points are ordered feature pairs that meet criteria for m/z, RT (rty vs rtProj), and Q tolerances set in the mzfit list argument (see: mzfitParam). If

Setting the plot option to TRUE generates an image of the curve fit through the selected points and is a useful method for determining if m/z mapping is appropriate for the analysis and tuning certain parameters.

This function is called within calcScores, which can help improve the pairwise scores

54 mzfitParam

Value

model object of class gam or 0 (if no model selected)

Examples

mzfitParam

Parameterize m/z Shift Model

Description

Return a list of parameters for computing a m/z shift model between a pair of LC-MS metabolomics data sets.

Usage

```
mzfitParam(mz = 0.003, rt = 0.03, Q = 0.3, k = 10)
```

Arguments

mz	Numeric. Limits the m/z distance between feature pairs for modeling
rt	Numeric. Limits the RT distance between feature pairs for modeling (note: this is proportion of total retention time)
Q	Numeric. Limits the Q distance between feature pairs for modeling
k	integer k value controlling the dimension of the basis spline fit

Details

Correcting for systematic m/z shifts improves the scores for feature pair alignments, yielding more accurate match hypotheses. This function yields a parameter list for GAM spline fitting of points that meet criteria for m/z, RT (rty vs rtProj), and Q tolerances. The number of knots for the GAM fit is controlled by hyperparameter k.

mzGroup 55

Value

list of parameter values

Examples

```
mzfit \leftarrow mzfitParam(mz = 0.003, rt = 0.05, Q = 0.2, k = 20)
```

mzGroup

Binning of mass spectral features in m/z dimension

Description

Features in two input feature lists are grouped by their m/z values.

Usage

```
mzGroup(xset, yset, binGap)
```

Arguments

xset data frame containing metabolomics features
 yset data frame containing metabolomics features
 binGap numeric gap value between consecutive sorted & pooled feature m/z values.

Details

The m/z values from both datasets are pooled, sorted, and binned by the binGap argument. Feature groups form when there is at least one pair of features from both datasets whose consecutive difference is less than binGap. Grouped features are joined together in combinedTable data report.

Value

list object containing updated xset & yset with group information

56 nonmatched

nonmatched

Get Nonmatched Features

Description

Features that lack a any counterparts in the complementary dataset may be obtained from this method. If data is set to "x" or "y", will retrieve data from the current X or Y dataset, respectively. If data is set to NULL, will retrieve the list of nonmatched features.

Usage

```
nonmatched(object, data = "x")
## S4 method for signature 'metabCombiner'
nonmatched(object, data = "x")
```

Arguments

object metabCombiner object

dataset identifier for metabCombiner objects; if NULL, returns full list of non-

matched features

Value

Data frame of non-matched features corresponding to data argument

Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.005)

nnmx <- nonmatched(p.comb, data = "x")
nnmy <- nonmatched(p.comb, data = "y")</pre>
```

objective 57

objective

Weight Parameter Objective Function

Description

This function evaluates the A, B, C weight parameters in terms of score separability of matching versus mismatching compound alignments. Higher objective function value imply a superior weight parameter selection.

Usage

```
objective(
  cTable,
  idtable,
 Α,
 В,
 С,
 minScore,
 mzdiff,
  rtdiff,
  qdiff,
  rtrange,
  adductdiff,
  penalty,
 matches,
  mismatches
)
```

Arguments

cTable	data frame. Abridged metabCombiner report table.
idtable	data frame containing all evaluated identities
A	Numeric weight for penalizing m/z differences.
В	Numeric weight for penalizing differences between fitted & observed retention times
С	Numeric weight for differences in Q (abundance quantiles).
minScore	numeric. Minimum score to count towards objective value.
mzdiff	numeric differences between feature m/z values
rtdiff	Differences between model-projected retention time value $\&$ observed retention time
qdiff	Difference between feature quantile Q values.
rtrange	range of dataset Y retention times
adductdiff	Numeric divisors of computed score when non-empty adduct labels do not match
penalty	positive numeric penalty wherever $S(i,j) > S(i,i)$, $i = -j$

58 opts.duplicate

matches integer row indices of identity matches

mismatches list of integer identity row mismatches for each identity

Details

First, the similarity scores between all grouped features are calculated as described in scorePairs Then, the objective value for a similarity S is evaluated as:

$$OBJ(S) = \sum h(S(i,i)) - h(S(i,j)) - p(S(i,i) > S(i,j))$$

- -S(i,i) represents the similarity between correct identity alignments
- -S(i,j), represents the maximum similarity of i to grouped feature j, i =/= j (the highest-scoring misalignment)
- -h(x) = x if x > minScore, 0 otherwise
- -p(COND) = 0 if the condition is true, and a penalty value otherwise

This is summed over all labeled compound identities (e.g. idx = idy) shared between input datasets.

Value

A numeric value quantifying total separability of compound match similarity scores from mismatch scores, given A,B,C values

opts.duplicate

Duplicate Feature Detection Parameters

Description

Lists the parameters for detection of two or more rows that represent the same entity, based on similar m/z and retention time values.

Usage

```
opts.duplicate(
  mz = 0.0025,
  rt = 0.05,
  resolve = c("single", "merge"),
  weighted = FALSE
)
```

Arguments

mz m/z tolerance for duplicate feature detection

rt RT tolerance for duplicate feature

resolve character. Either "single" (default) or "merge".

weighted logical. Option to weight m/z, RT, Q by mean abundance of each row (TRUE)

or take single representative values (FALSE).

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Details

The presence of duplicate features has negative consequences for the LC-MS alignment task. The package offers several options for resolving the issue of feature duplication. Pairwise m/z and RT tolerances define which features are to be considered as duplicates within a single data set. Setting mz or rt to 0 skips duplicate feature filtering altogether.

When duplicates are detected, either a single master copy is retained (resolve = "single") or merged into a single row (resolve = "merge"). The master copy is the copy with lower proportion of missingness, followed by the most abundant (by median or mean). If missingness and abundance is equivalent for duplicates, the first copy that appears is retained. The "merge" option fuses duplicate feature rows, with quantitative descriptors (m/z, RT) either calculated as a weighted average (weighted = TRUE) or otherwise taken from the top representative row; id and adduct values are concatenated; the maximum feature value is used for each sample; and all 'extra' values are taken from the 'master copy' row, similar to the "single" option.

Examples

```
data(plasma20)
pars.duplicate <- opts.duplicate(mz = 0.01, rt = 0.05, resolve = "single")
p20 <- metabData(plasma20, samples = "Red", duplicate = pars.duplicate)

#to prevent removal of duplicate features
p20 <- metabData(plasma20, samples = "Red", duplicate = opts.duplicate(0))

##merge option
pars.duplicate <- opts.duplicate(mz = 0.01, rt = 0.05, resolve = "merge")
p20 <- metabData(plasma20, samples = "Red", duplicate = pars.duplicate)</pre>
```

plasma20

20 minute LC-MS Analysis of Human Plasma

Description

An example metabolomics analysis of human plasma from Red Cross and CHEAR cohorts, plus pooled aliquots and blanks, acquired with a 20 minute total Reversed-Phase Liquid Chromatography & QTOF-MS instrument in the positive ionization mode.

Usage

```
data(plasma20)
```

Format

A data frame with 8910 rows and 22 columns.

plasma30

30 minute LC-MS Analysis of Human Plasma

Description

An example metabolomics analysis of human plasma from Red Cross and CHEAR cohorts, plus pooled aliquots and blanks, acquired with a 30 minute total Reversed-Phase Liquid Chromatography and a QTOF-MS instrument in the positive ionization mode.

Usage

```
data(plasma30)
```

Format

A data frame with 8286 rows and 22 columns

```
{\tt plot,metabCombiner,ANY-method} \\ {\tt Plot\ metabCombiner\ Fits}
```

Description

This is a plotting method for metabCombiner objects. It displays ordered pairs and a curve fit computed using fit_gam or fit_loess, using base R graphics.

Usage

```
## S4 method for signature 'metabCombiner,ANY'
plot(x, y, ...)

plot_fit(
   object,
   fit = c("gam", "loess"),
   pcol = "black",
   lcol = "red",
   lwd = 3,
   pch = 19,
   outlier = "show",
   ocol = "springgreen4",
   legend = c("anchor", "outlier"),
   ...
)
```

Arguments

x	metabCombiner object
У	
	Other variables passed into graphics::plot
object	metabCombiner object
fit	choice of model (either "gam" or "loess").
pcol	color of the normal points (ordered RT pair) in the plot
lcol	color of the fitted line in the plot
lwd	line width of the curve fit between anchor points
pch	plot character type; see ?graphics::par for details
outlier	display option for outliers. If "show" or "s", treats outlier points like normal anchors; if "remove" or "r", removes outlier points from the plot; if "highlight" or "h", displays outliers with a different color and associated legend.
ocol	color of the outlier points; outlier argument must be set to "highlight" or "h"
legend	length-2 character vector indicating point labels in the legend if outlier argument set to "highlight" or "h"

Value

no values returned

Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb = metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)
p.comb = selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
p.comb = fit_gam(p.comb, k = 20, iterFilter = 1, family = "gaussian")

##plot of GAM fit
plot(p.comb, main = "Example GAM Fit Plot", xlab = "X Dataset RTs",
    ylab = "Y Dataset RTs", pcol = "red", lcol = "blue", lwd = 5,
    fit = "gam", outliers = "remove")

grid(lwd = 2, lty = 3) #adding gridlines</pre>
```

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QData

Retrieve Relative Abundance Values

Description

This retrieves feature Q values from one or all constituent dataset features of a metabCombiner object. Alternatively, the average Q value can be retrieved.

Usage

```
QData(object, data = NULL, value = c("obs", "mean"))
## S4 method for signature 'metabCombiner'
QData(object, data = NULL, value = c("observed", "mean"))
```

Arguments

object metabCombiner object

data dataset identifier to extract information from; if NULL, extracts information

from all datasets

value Either "obs" (observed - default option) or "mean" average value

Value

```
data frame or vector of relative ranked abundance (Q) values
data(plasma30) data(plasma20)
p30 <- metabData(head(plasma30,500), samples = "CHEAR") p20 <- metabData(head(plasma20,500),
samples = "Red") p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")
##retrieve all Q Q <- QData(p.comb, data = NULL)
##retrieve Q from p30 Q <- QData(p.comb, data = "p30")
##retrieve mean Q Q <- QData(p.comb, value = "mean")
```

resolveRows

Resolve Conflicting Alignment Subgroups

Description

This method resolves conflicting feature pair assignments (labeled as "CONFLICT") to obtain 1-1 feature matches in the combinedTable results report.

Usage

```
resolveRows(fields, rtOrder)
```

rtData 63

Arguments

fields data frame containing the main

rtOrder logical option to impose RT order for resolving subgroups

Details

This is called from within labelRows (with argument resolveConflicts set to TRUE), reduceTable, & metabCombiner (using metabCombiner object inputs). The method determines which combination of unique feature pairs has the highest sum of scores ("resolveScore") within each subgroup. By default, these combinations of feature pairs must have consistency in their retention time order (rtOrder = TRUE). The combination of 1-1 feature pair alignments with the highest resolveScore within the subgroups are annotated as "RESOLVED", with the remaining unannotated rows labeled as "REMOVE" (or removed outright by other package functions). Feature pairs belonging to multiple subgroup (alt > 0) are labeled as REMOVE.

Value

data.frame of combinedTable fields, replacing "CONFLICT" labels with "RESOLVED" or "RE-MOVE", depending on the computations performed.

rtData Retrieve Retention Time Values

Description

This retrieves feature RT values from one or all constituent dataset features of a metabCombiner object. Alternatively, the average RT value can be retrieved.

Usage

```
rtData(object, data = NULL, value = c("obs", "mean"))
## S4 method for signature 'metabCombiner'
rtData(object, data = NULL, value = c("observed", "mean"))
```

Arguments

object metabCombiner object

dataset identifier to extract information from; if NULL, extracts information

from all datasets

value Either"obs" (observed - default option) or "mean"

Value

data frame or vector of retention time values

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Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(head(plasma30,500), samples = "CHEAR")
p20 <- metabData(head(plasma20,500), samples = "Red")
p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")

##retrieve all RTs
rt <- rtData(p.comb, data = NULL)

##retrieve RTs from p30
rt <- rtData(p.comb, data = "p30")

##retrieve mean RT
rt <- rtData(p.comb, value = "mean")</pre>
```

scorePairs

Calculate Pairwise Alignment Scores

Description

Helper function for calcScores & evaluateParams. Calculates a pairwise similarity score between grouped features using differences in m/z, rt, and Q.

Usage

```
scorePairs(A, B, C, mzdiff, rtdiff, qdiff, rtrange, adductdiff)
```

Arguments

Α	Numeric weight for penalizing m/z differences.
В	Numeric weight for penalizing differences between fitted & observed retention times.
С	Numeric weight for differences in Q (abundance quantiles).
mzdiff	Numeric differences between feature m/z values
rtdiff	Differences between model-projected retention time value $\&$ observed retention time
qdiff	Difference between feature quantile Q values
rtrange	Range of dataset Y retention times
adductdiff	Numeric divisors of computed score when non-empty adduct labels do not match

selectAnchors 65

Details

The score between two grouped features x & y is calculated as:

```
S = -exp(-A|mzx - mzy| - B|rty - rtproj|/rtrange - C|Qx - Qy|)
```

where mzx & Qx correspond to the m/z and abundance quantile values of feature x; mzy, rty, and Qy correspond to the m/z, retention time, and quantile values of feature y; rtproj is the model-projected retention time of feature x onto the Y dataset chromatogram and rtrange is the retention time range of the Y dataset chromatogram. A, B, C are non-negative constant weight parameters for penalizing m/z, rt, and Q differences. Values between 0 (no confidence alignment) and 1 (high confidence alignment).

Value

Numeric similarity score between 0 & 1

selectAnchors

Select Anchors for Nonlinear RT Model

Description

A subset of possible alignments in the combinedTable are used as ordered pairs to anchor a retention time projection model. Alignments of abundant features are prominent targets for anchor selection, but shared identified features (i.e. feature pairs where idx = idy) may be used.

Usage

```
selectAnchors(
  object,
  useID = FALSE,
  tolmz = 0.003,
  tolQ = 0.3,
  tolrtq = 0.3,
  windx = 0.03,
  windy = 0.03,
  brackets_ignore = c("(", "[", "{"}")")
```

Arguments

```
object metabCombiner object.

useID logical. Option to first search for IDs as anchors.

tolmz numeric. m/z tolerance for prospective anchors

tolQ numeric. Quantile Q tolerance for prospective anchors

tolrtq numeric. Linear RT quantile tolerance for prospective anchors.
```

66 selectAnchors

windx numeric. Retention time exclusion window around each anchor in X dataset.

Optimal values are between 0.01 and 0.05 min (1-3s)

windy numeric. Retention time exclusion window around each anchor in dataset Y.

Optimal values are between 0.01 and 0.05 min (1-3s)

brackets_ignore

If useID = TRUE, bracketed identity strings of the types included in this argu-

ment will be ignored.

Details

In order to map between two sets of retention times, a set of ordered pairs need to be selected for the spline fit. This function relies on mutually abundant features to select these ordered pairs. In iterative steps, the most abundant (as indicated by Q value) in one dataset is selected along with its counterpart, and all features within some retention time window specified by windx & windy arguments are excluded. This process is repeated until all features have been considered.

tolQ & tolmz arguments restrict to feature pairs that have differences in Q & m/z within these tolerances. tolrtq further limits to feature pairs those with relative differences in linear retention time quantiles, calculated as rtqx = (rtx - min(rtx))/(max(rtx) - min(rtx)) & rtqy = (rty - min(rty))/(max(rty) - min(rty))

Shared identities (in which idx & idy columns have matching, non-empty & non-bracketed strings) may be used if useID is set to TRUE. In this case, shared identities will be searched first and will not be subject to any of the restrictions in m/z, Q, or rt. The iterative process proceeds after processing of shared identities.

Value

labels

metabCombiner object with updated anchors slot. This is a data.frame of feature pairs that shall be used to map between retention times using a GAM or LOESS model.

anchor labels; "I" for identity, "A" for normal anchors

idx	identities of features from dataset X
idy	identities of features from dataset Y
mzx	m/z values of features from dataset X
mzy	m/z values of features from dataset Y
rtx	retention time values of features from dataset X
rty	retention time values of features from dataset Y
rtProj	model-projected retention time values from \boldsymbol{X} to \boldsymbol{Y}
Qx	abundance quantile values of features from dataset \boldsymbol{X}
Qy	abundance quantile values of features from dataset Y
adductX	adduct label of features from dataset X
adductY	adduct label of features from dataset Y
group	m/z feature group of feature pairing

selectAnchorsParam 67

See Also

```
getAnchors, fit_gam, fit_loess
```

Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.005)

##example 1 (no known IDs used)
p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windx = 0.03, windy = 0.02, tolrtq = 0.3)

##example 2 (known IDs used)
p.comb <- selectAnchors(p.comb, useID = TRUE, tolmz = 0.003, tolQ = 0.3)

##To View Plot of Ordered Pairs
anchors = getAnchors(p.comb)
plot(anchors$rtx, anchors$rty, main = "Selected Anchor Ordered Pairs", xlab = "rtx", ylab = "rty")</pre>
```

selectAnchorsParam

List selectAnchors Defaults

Description

List of default parameters for anchor selection step of main package workflow, which can be used as input for the wrapper functions. See help(selectAnchors) or ?selectAnchors for more details.

Usage

```
selectAnchorsParam(
  useID = FALSE,
  tolmz = 0.003,
  tolQ = 0.3,
  tolrtq = 0.3,
  windx = 0.03,
  windy = 0.03,
  brackets_ignore = c("(", "[", "{"}")")
```

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Arguments

useID	Choice of using IDs for anchor selection; default: FALSE
tolmz	m/z tolerance for ordered pair features; default: 0.003
tolQ	Q tolerance for ordered pair features; default: 0.3

tolrtq RT quantile tolerance for ordered pair features; default: 0.5

windx X feature RT window parameter. Default: 0.03 windy Y feature RT window parameter. Default: 0.03

brackets_ignore

bracket types for ignoring string comparisons

Value

list of selectAnchors parameters

See Also

selectAnchors, metabCombine

Examples

```
sa_param <- selectAnchorsParam(tolmz = 0.002, tolQ = 0.2, windy = 0.02)</pre>
```

updateTables Update metabCombiner Objects

Description

This method updates the feature list (featData) and aligned table (combinedTable) within a metabCombiner object. Manual changes to the (combinedTable) as well as unmatched X & Y dataset features can be incorporated into the object and the corresponding results. This function is typically paired with link{reduceTable} or other forms of table reduction performed by the user.

Usage

```
updateTables(object, xdata = NULL, ydata = NULL, combinedTable = NULL)
```

Arguments

object	metabCombiner	object to	be updated
--------	---------------	-----------	------------

xdata metabData or metabCombiner object originally used to construct the object ar-

gument

ydata metabData or metabCombiner object originally used to construct the object ar-

gument

combinedTable merged table which may be altered by the user. This must have the combinedTable

format to be valid (see: ?isCombinedTable)

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Details

There are two points where features can be removed from the combinedTable report: during m/z grouping and during the table reduction step. It is also possible for user-specified changes to the report to remove certain features entirely. This function allows for the missed features to be brought back into the table as non-matched entities. For xdata features, the Y columns will be entirely missing values, and ydata features will have missing X information. The feature data (featData) will also be updated for use in subsequent alignments, but only features present in the representative dataset will be retained by default.

Value

metabCombiner object with updates to combinedTable to include features that have been missed or changes by the user.

Note

Duplicated sample & extra column names cannot be copied from the original data they feature in, therefore they are left as missing values.

Examples

```
data(plasma30)
data(plasma20)
p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red")</pre>
p.comb <- metabCombiner(xdata = p30, ydata = p20, xid = "p30", yid = "p20",
                         binGap = 0.0075)
##extracting, modifying, and updating combinedTable
cTable <- combinedTable(p.comb)</pre>
cTable <- dplyr::filter(cTable, rty < 17.25)
p.comb <- updateTables(p.comb, combinedTable = cTable)</pre>
p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolQ = 0.3, windy = 0.02)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 1)</pre>
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.5)
p.comb <- reduceTable(p.comb, delta = 0.2, maxRTerr = 0.5)</pre>
##updating to include features removed from xdata & ydata
p.comb <- updateTables(p.comb, xdata = p30, ydata = p20)</pre>
#view results
cTable <- combinedTable(p.comb)</pre>
fdata <- featData(p.comb)</pre>
```

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write2file

Print metabCombiner Report to File.

Description

Prints a combinedTable report to a file, specified by file argument. Output file has an empty line between each separate m/z group for ease of viewing.

Usage

```
write2file(object, file, sep = ",")
```

Arguments

object metabCombiner object or combinedTable

file character string naming the output file path

sep Character field separator. Values within each row are separated by this character.

Value

no values returned

Examples

```
data(plasma30)
data(plasma20)

p30 <- metabData(plasma30, samples = "CHEAR")
p20 <- metabData(plasma20, samples = "Red", rtmax = 17.25)
p.comb <- metabCombiner(xdata = p30, ydata = p20, binGap = 0.0075)

p.comb <- selectAnchors(p.comb, tolmz = 0.003, tolrtq = 0.3, windy = 0.02)
p.comb <- fit_gam(p.comb, k = 20, iterFilter = 1)
p.comb <- calcScores(p.comb, A = 90, B = 14, C = 0.5)
p.comb <- labelRows(p.comb, maxRankX = 2, maxRankY = 2, remove = TRUE)

###using metabCombiner object as input
write2file(p.comb, file = "plasma-combined.csv", sep = ",")

###using combinedTable report and feature data as input
cTable <- combinedTable(p.comb)
write2file(cTable, file = "plasma-combined.txt", sep = "\t")</pre>
```

x 71

Obtain x & y Data Identifiers

Χ

Description

metabCombiner alignment is performed in a pairwise manner between two datasets generically termed "x" & "y". These methods print the identifier(s) associated with datasets X and Y, contained within the xy slot of a constructed metabCombiner object.

Usage

```
x(object)
xy(object)

## S4 method for signature 'metabCombiner'
x(object)

## S4 method for signature 'metabCombiner'
xy(object)

## S4 method for signature 'metabCombiner'
y(object)
```

Arguments

object metabCombiner object

Value

```
character X or Y dataset identifiers
data(plasma30) data(plasma20)
p30 <- metabData(head(plasma30,500), samples = "CHEAR") p20 <- metabData(head(plasma20,500), samples = "Red") p.comb <- metabCombiner(p30, p20, xid = "p30", yid = "p20")
#expected: "p30" x(p.comb)
#expected: "p20" y(p.comb)
#list of x & y data descriptors xy(p.comb)
```

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