

Package ‘MetabolomicsBasics’

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Type Package

Title Basic Functions to Investigate Metabolomics Data Matrices

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Description A set of functions to investigate raw data from (metabol)omics experiments intended to be used on a raw data matrix, i.e. following peak picking and signal deconvolution. Functions can be used to normalize data, detect biomarkers and perform sample classification. A detailed description of best practice usage may be found in the publication <[doi:10.1007/978-1-4939-7819-9_20](https://doi.org/10.1007/978-1-4939-7819-9_20)>.

URL <https://github.com/janlisec/MetabolomicsBasics>

License GPL-3

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AdjustSymbols	<i>AdjustSymbols.</i>
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Description

AdjustSymbols will generate plotting character and color vectors based on experimental factors.#'

Usage

AdjustSymbols(cols = NULL, pchs = NULL, colorset = NULL, symbolset = NULL)

Arguments

cols	Factor (color output) or numeric (grey-scale output) vector or NULL (omitted).
pchs	Factor vector or NULL (omitted).
colorset	Color definitions for the factor levels of 'cols' (can be omitted to use default values).
symbolset	Plotting character definitions for the factor levels of 'pchs' (can be omitted to use default values).

Details

Using a fixed color and symbol scheme indicating samples from different groups throughout all figures of a analysis workflow is a reasonable decision. This function allows to specify both and attach it to a sample table for further use.

Value

Either a vector (if one parameter of ‘cols’ and ‘pchs’ remains NULL), a data frame with columns ‘cols’ and ‘pchs’ (if both are provided and of equal length) or a list of length 2 (if both are provided and of different length). Will be used by several plotting functions of the package internally.

Examples

```
# return color vector
x <- gl(6, 3)
y <- as.numeric(x)
plot(y, bg = AdjustSymbols(cols = x), pch = 21, cex = 2)
plot(y, bg = AdjustSymbols(cols = y), pch = 21, cex = 2)
plot(y, bg = AdjustSymbols(cols = x, colorset = 1:6), pch = 21, cex = 2)
plot(y, pch = AdjustSymbols(pchs = x), cex = 2)
plot(y, bg = 2, pch = AdjustSymbols(pchs = x, symbolset = 1:6), cex = 2)

# load data and plot using provided color scheme
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam
head(sam)
plot(y = raw[, 1], x = as.numeric(sam$GT), pch = sam$pchs, bg = sam$cols)

# change colors to greyscale
head(AdjustSymbols(cols = sam$GT, pchs = sam$Origin))
tmp.set <- grDevices::rainbow(length(levels(sam$GT)))
head(AdjustSymbols(cols = sam$GT, pchs = sam$Batch, colorset = tmp.set))
plot(raw[, 1] ~ sam$GT, col = unique_labels(sam = sam, g = "GT")[, "cols"])
sam$cols <- AdjustSymbols(cols = as.numeric(sam$GT))
plot(raw[, 1] ~ sam$GT, col = unique_labels(sam = sam, g = "GT")[, "cols"]) #'
```

CheckForOutliers

CheckForOutliers.

Description

CheckForOutliers will evaluate a numeric vector and check if outliers within groups based on group $\text{mean} \pm n \times \text{sd}$.

Usage

```
CheckForOutliers(
  x = NULL,
  group = NULL,
  n_sd = 3,
  method = c("idx", "logical", "dist")
)
```

Arguments

x	Numeric vector.
group	Factor vector of length(x).
n_sd	Cutoff for outliers in E being $\text{mean}(\text{group}) + n_sd * \text{sd}(\text{group})$ where group values are calculated without the outlier candidate.
method	Different variants of the result value. See details.

Details

The numeric will be split by groups and each value will be evaluated with respect to its distance to the group mean (calculated out of the other values in the group). Distance here means the number of standard deviations the value is off the group mean. With different choices of method the output can be switched from the calculated fold-distances to a boolean of length(x) or and Index vector giving the outliers directly (see examples).

Value

Depending on the selected method. See details.

Examples

```

set.seed(0)
x <- runif(10)
x[1] <- 2
group <- gl(2, 5)
CheckForOutliers(x, group, method = "dist")
CheckForOutliers(x, group, method = "logical")
CheckForOutliers(x, group, method = "idx")
graphics::par(mfrow = c(1, 2))
bg <- c(3, 2)[1 + CheckForOutliers(x, group, method = "logical")]
graphics::plot(x = as.numeric(group), y = x, pch = 21, cex = 3,
  bg = bg, main = "n_sd=3", las = 1, xlim = c(0.5, 2.5))
bg <- c(3, 2)[1 + CheckForOutliers(x, group, n_sd = 4, method = "logical")]
graphics::plot(x = as.numeric(group), y = x, pch = 21, cex = 3,
  bg = bg, main = "n_sd=4", las = 1, xlim = c(0.5, 2.5))
graphics::par(mfrow = c(1, 1))

# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam

# no missing data in this matrix
all(is.finite(raw))

# check for outliers (computing n-fold sd distance from group mean)
tmp <- apply(raw, 2, CheckForOutliers, group = sam$GT, method = "dist")
# plot a histogram of the observed distances
graphics::hist(tmp, breaks = seq(0, ceiling(max(tmp))), main = "n*SD from mean", xlab = "n")

# Calculate the amount of values exceeding five-sigma and compare with a standard gaussian

```

```

table(tmp > 5)
round(100 * sum(tmp > 5) / length(tmp), 2)

gauss <- CheckForOutliers(x = rnorm(prod(dim(raw))), method = "dist")
sapply(1:5, function(i) {
  data.frame("obs" = sum(tmp > i), "gauss" = sum(gauss > i))
})

# compare a PCA w/wo outliers
RestrictedPCA(
  dat = raw, sam = sam, use.sam = sam$GT %in% c("Mo17", "B73"), group.col = "GT",
  fmod = "GT+Batch+Order", P = 1, sign.col = "GT", legend.x = NULL, text.col = "Batch", medsd = TRUE
)
raw_filt <- raw
raw_filt[tmp > 3] <- NA
RestrictedPCA(
  dat = raw_filt, sam = sam, use.sam = sam$GT %in% c("Mo17", "B73"), group.col = "GT",
  fmod = "GT+Batch+Order", P = 1, sign.col = "GT", legend.x = NULL, text.col = "Batch", medsd = TRUE
)

```

ClassificationCV

ClassificationCV.

Description

ClassificationCV will perform a classification using SVM's and/or Decision Trees including cross validation on a data set according to a provided grouping vector.

Usage

```

ClassificationCV(
  d = NULL,
  g = NULL,
  n = 1,
  k = 1,
  rand = F,
  method = c("svm", "C50", "rpart", "ropls"),
  method.control = list(),
  silent = FALSE
)

```

Arguments

d	Data matrix or data.frame with named rows (samples) and columns (traits).
g	Group-vector, factor.
n	Replicates of classifications.
k	Number of folds per replicate.

rand	Randomize Group-vector (and apply according n and k to this randomization).
method	Currently svm, ropls and decision tree methods C50 and rpart are supported.
method.control	A list of parameters, forwarded to the respective classification function.
silent	Logical. Set TRUE to suppress progress bar and warnings.

Details

This function allows to demonstrate the functionality of different classification tools with respect to building classifiers for metabolomics data. Check the examples in [ClassificationWrapper](#) for automatic multi-fold analysis.

Value

A list of classification results which can be analyzed for accuracy, miss-classified samples and more.

ClassificationHistogram

ClassificationHistogram.

Description

ClassificationHistogram will plot the results of [ClassificationWrapper](#).

Usage

```
ClassificationHistogram(out_classific = NULL, breaks = seq(0, 1, 0.05), ...)
```

Arguments

out_classific	Output of ClassificationWrapper .
breaks	Breaks for histogram.
...	Passed on to par. Useful to adjust cex.

Details

No further details.

Value

Returns NULL invisibly.

ClassificationWrapper *ClassificationWrapper*.

Description

ClassificationWrapper will do classification using SVM's and/or Decision Trees including cross validation.

Usage

```
ClassificationWrapper(
  d = NULL,
  g = NULL,
  n = 100,
  n_rand = 1,
  k = 5,
  method = c("C50", "svm", "rpart", "ropls"),
  train = NULL,
  method.control = list(),
  silent = FALSE
)
```

Arguments

d	data, matrix or data.frame !! needs row/col-names.
g	Group-vector, factor.
n	replicates of classifications, i.e. number of different split into folds.
n_rand	different number of randomizations, see Details.
k	Fold cross validation.
method	Currently svm, ropls and decision tree methods (C50 and rpart) are supported.
train	Either NULL (random permutations) or an index vector for a training subset out of g.
method.control	A list of parameters, forwarded to the selected methods function.
silent	Logical. Set TRUE to suppress progress bar and warnings.

Details

Parameter 'n_rand' will influence how permutation testing for robustness is conducted. If n_rand=1 than samples will be permuted exactly one time and subjected to n replications (with respect to fold splitting). If n_rand>1, samples will be permuted this many times but number of replications will be lowered to limit processing time. A good compromise is to balance both, using less replications than for observed data but on several randomizations.

Value

#' Classification results as list.

Examples

```
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam
gr <- sam$Origin
# establish a basic rpart model and render a fancy plot including the accuracy

class_res <- ClassificationWrapper(d = raw, g = gr, method = c("rpart", "svm"), n = 3, k = 3)
ClassificationHistogram(class_res)
```

find_boundaries	<i>find_boundaries.</i>
-----------------	-------------------------

Description

find_boundaries will determine peak boundaries within a BPC or mass trace.

Usage

```
find_boundaries(
  int = NULL,
  rt = NULL,
  p = which.max(int),
  k = 3,
  bl = min(int),
  local_min = int[p]
)
```

Arguments

int	The measured intensity of the ion mass (obviously ordered according to consecutive RTs).
rt	The respective retention times (can be omitted as currently not used).
p	The anticipated peak position (as index of int) if several peaks are within the mass trace.
k	The smoothing window parameter (provided to runmed).
bl	The baseline value. Can be provided explicitly if automatic determination is insufficient.
local_min	This is practically the upper end of the baseline. It can be set to avoid boundary detection at local minima (e.g. for peaks suffering ion suppression).

Details

It is yet another peak finder or, more precisely, it is a function to identify two RT values which flank a intensity maximum which is required if one would like to integrate the peak area.

Value

Numeric vector of length=2 specifying the start and end index of the peak.

Examples

```
int <- sin(seq(-0.75 * pi, 1.75 * pi, by = 0.1))
plot(int)
abline(v = find_boundaries(int = int))
```

MBoxplot

MBoxplot.

Description

MBoxplot will generate an annotated boxplot. A unifying function for MS-data Boxplots based on 'raw' and 'sam'.

Usage

```
MBoxplot(
  pk = pk,
  raw = NULL,
  sam = NULL,
  met = NULL,
  g = NULL,
  flt = NULL,
  an = NULL,
  plot_sample_n = FALSE,
  txt = NULL,
  cex.txt = 0.5,
  plot_rel_axis = NULL,
  ...
)
```

Arguments

pk	Colname of raw to plot if pk is character OR the colnum number if pk is numeric.
raw	Plotting data as samples (rows) x metabolites (cols).
sam	Sample table.
met	Containing at minimum columns for annotation (see parameter an) and nrow(met) should be ncol(raw).
g	Grouping vector if Group not contained in sam.
flt	Filter to exclude certain samples (T/F) vector.
an	Switch to include annotation (from met) in the boxplot providing a character vector of colnames from met.

<code>plot_sample_n</code>	Amend each box with the number of finite values which were a basis for plotting this group.
<code>txt</code>	Character vector with information per sample to be plotted on top of the box as text.
<code>cex.txt</code>	Specify size of annotation text.
<code>plot_rel_axis</code>	Specify one level of <code>g</code> (or <code>sam\$Group</code>) which to express the data relative against.
<code>...</code>	Further options parsed to <code>boxplot</code> .

Details

Usually metabolomics experiments are conducted on multiple replicates of a sample group. Box-plots allow to quickly access potential differences between measurement values of several groups. `MBoxplot` can be nicely used to generate QC plots for all metabolites prior and after normalization, in absolute or relative scale and sorted according to significance.

Value

Nothing. Will produce a plot (or file if specified).

Examples

```
x <- data.frame("y" = runif(36), "GT" = gl(3, 12), "TP" = factor(rep(rep(1:3, each = 4), 3)))
x <- cbind(x, AdjustSymbols(cols = x$GT, pchs = x$TP))
MBoxplot(
  pk = "y", raw = x, sam = x, met = data.frame("Peak" = "y", "Test" = I("info")),
  g = interaction(x$GT, x$TP), an = "Test", plot_n_samples = TRUE, txt = rownames(x)
)
```

met

Metabolite table

Description

This data frame contains the metabolite definition of 112 metabolites according to the cols of [raw](#).

Usage

```
met
```

Format

An object of class `data.frame` with 112 rows and 2 columns.

Author(s)

Jan Lisec <jan.lisec@bam.de>

Source

<<https://onlinelibrary.wiley.com/doi/10.1111/j.1365-313X.2011.04689.x>>

MetaboliteANOVA

MetaboliteANOVA.

Description

MetaboliteANOVA will perform an ANOVA on columns of a data matrix according to a specified model.

Usage

```
MetaboliteANOVA(  
  dat = NULL,  
  sam = NULL,  
  model = NULL,  
  method = "none",  
  silent = FALSE  
)
```

Arguments

<code>dat</code>	Data matrix (e.g. of metabolite).
<code>sam</code>	Sample table (same number of row as 'dat' and containing all columns specified in 'model').
<code>model</code>	ANOVA model. May include +, * and : together with column names of sam (cf. Examples).
<code>method</code>	The method to be used in column wise multiple testing adjustment, see p.adjust .
<code>silent</code>	Logical. Shall the function print warnings to the console?

Details

The function is a wrapper for `lm` including some sanity checks. It will accept a data matrix (traits in columns), sample information (data.frame) and a potential model as input, compute an ANOVA per column and return the respective P-values in a named matrix for further plotting or export.

Value

A named matrix of P-values (rows=metabolites/traits; cols=ANOVA factors).

Examples

```
# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam
# compute P-values according to specified ANOVA model (simple and complex)
head(m1 <- MetaboliteANOVA(dat = raw, sam = sam, model = "GT"))
head(m2 <- MetaboliteANOVA(dat = raw, sam = sam, model = "GT+Batch+Order+MP"))
# compare P-values for one factor determined in both models
hist(log10(m2[, "GT"]) - log10(m1[, "GT"]), main = "")
```

msconvert

msconvert.

Description

msconvert is calling ProteoWizards MSConvert as a command line tool on Windows.

Usage

```
msconvert(
  files = NULL,
  msc_exe =
    "C:\\Program Files\\ProteoWizard\\ProteoWizard 3.0.11856\\msconvert.exe",
  args = c("--filter \"peakPicking cwt snr=0.01 peakSpace=0.1 msLevel=1\"",
    "--filter \"scanTime [0,3600]\"", "--filter \"metadataFixer\"", "--mzML", "--32",
    "--zlib")
)
```

Arguments

files	A character vector of MS data files (wiff, raw, d, ...).
msc_exe	The path to the installed 'msconvert.exe'.
args	The arguments passed to msconvert on the command line (see details for documentation).

Details

It is a quick and dodgy function to show how to convert vendor MS data into an open format (mzML). You will have to download/install MSConvert prior to usage, and probably adjust the arguments according to your needs. Arguments are documented here <https://proteowizard.sourceforge.io/tools/msconvert.html>. If you don't know where the msconvert.exe is installed you can check for the correct path using `list.files(path="C:/", pattern="^msconvert.exe$", recursive = TRUE)`.

Value

Only some informative outputs printed to the console. The specified MS data files will be converted to mzML within the same folder.

PlotMetabolitePCA	<i>PlotMetabolitePCA.</i>
-------------------	---------------------------

Description

PlotMetabolitePCA will show PC1 and PC2 of a `pcaMethods` object and generate a flexible plot.

Usage

```
PlotMetabolitePCA(  
  pca_res = NULL,  
  sam = NULL,  
  g = NULL,  
  medsd = FALSE,  
  text.col = "ID",  
  legend.x = "bottomleft",  
  comm = NULL  
)
```

Arguments

<code>pca_res</code>	A <code>pcaRes</code> object from the <code>pcaMethods</code> package.
<code>sam</code>	Sample table including columns 'cols', 'pchs' (for data point color and shape) and 'ID' (to label data points) 'Group' (to split cols for legend) 'MP' (to adjust point size).
<code>g</code>	Can be a factor vector of length= <code>nrow(sam)</code> and will influence legend and 'medsd'.
<code>medsd</code>	Calculate mean and sd for groups and overlay PCA plot with this information.
<code>text.col</code>	Data points may be overlaid by textual information, e.g. sample ID and 'text.col' specifies the column name of 'sam' to use for this purpose.
<code>legend.x</code>	Position of a legend or NULL to omit it.
<code>comm</code>	Will print commentary text to the bottom right of the plot (can be a character vector).

Details

See examples.

Value

A vector fo similar length as input but with various name components removed.

Examples

```
# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam

# calculate pca Result using pcaMethods and plot
pca_res <- pcaMethods::pca(raw, method = "rnipals", scale = c("none", "pareto", "uv")[2])
PlotMetabolitePCA(pca_res = pca_res, sam = sam, g = sam$GT)
# plot without legend and Group means instead
PlotMetabolitePCA(
  pca_res = pca_res, sam = sam, g = sam$GT, legend.x = NULL, text.col = NULL,
  medsd = TRUE, comm = LETTERS[1:4]
)

# readjust symbols before plotting
sam$Group <- interaction(sam$Origin, sam$Class, sep = "_")
sam[, c("cols", "pchs")] <- AdjustSymbols(cols = sam$Group, pchs = sam$Group)
PlotMetabolitePCA(pca_res = pca_res, sam = sam, g = sam$Group)
```

PlotPValueHist

PlotPValueHist.

Description

PlotPValueHist will take a named matrix of P-values (i.e. numeric between 0..1) and plot histograms for each column. In the easiest case this matrix is generated by [MetaboliteANOVA](#).

Usage

```
PlotPValueHist(
  out = NULL,
  method = "BH",
  x1 = "ANOVA P-values",
  y1 = "Number of metabolites",
  frac.col = NULL,
  ...
)
```

Arguments

out	matrix/data.frame; P-value table from 'MetaboliteANOVA.R' with factors in named columns and trait P-values in rows.
method	Multiple testing correction method applied, piped to p.adjust().
x1	xlab.
y1	ylab.
frac.col	Render histogram bars in stacked colors according to provided color vector (should be a vector of valid color names of length=nrow(out)).
...	Passed on to par. Useful to adjust cex.

Details

See examples.

Value

NULL. Will generate a P-value histogram plot.

Examples

```
# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam

# compute P-values according to specified ANOVA model (simple and complex)
head(pvals <- MetaboliteANOVA(dat = raw, sam = sam, model = "GT+Batch+Order"))
PlotPValueHist(out = pvals)

# adjust multiple testing correction method and y lable
PlotPValueHist(out = pvals, method = "none", y1 = "Number of Genes")

# color bars (by chance or according to a metabolite group)
PlotPValueHist(out = pvals, frac.col = rep(2:3, length.out = nrow(pvals)))
met <- MetabolomicsBasics::met
met$Name[grep("ine$", met$Name)]
PlotPValueHist(out = pvals, frac.col = 2 + 1:nrow(pvals) %in% grep("ine$", met$Name))
```

PolarCoordHeterPlot	<i>PolarCoordHeterPlot.</i>
---------------------	-----------------------------

Description

PolarCoordHeterPlot will draw a plot in polar coordinates visualizing heterosis effects according to a layout by Swanson-Wagner, where plot radius represents log2 of fold change between lowest and highest genotype and plot angle represents the ratio between lowest, intermediate and highest genotype.

Usage

```
PolarCoordHeterPlot(
  x,
  gt = c("P1", "P1xP2", "P2"),
  rev_log = NULL,
  exp_fac = 1,
  thr = 1,
  plot_lab = c("none", "text", "graph"),
  col = NULL
)
```

Arguments

<code>x</code>	Data matrix with measurement values (traits in rows and genotypes in columns).
<code>gt</code>	Character vector of length=3 indicating P1, F and P2. These are used to filter by column name from <code>x</code> .
<code>rev_log</code>	If you've log transformed your data, you might want to revert the log transformation.
<code>exp_fac</code>	Expansion factor to increase figure size.
<code>thr</code>	Alpha level used in ANOVA to filter insignificant rows. Keep <code>thr=1</code> to include all matrix rows.
<code>plot_lab</code>	Show 'text' or 'graph' style labels of the polar sections (or keep 'none' to omit).
<code>col</code>	Provide a color vector of length <code>nrow(x)</code> .

Details

See examples.

Value

Will generate a plot in polar coordinates and return the x/y coordinates of the data points invisibly.

Examples

```
# using the provided experimental data
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam
x <- t(raw)
colnames(x) <- sam$GT
gt <- c("B73", "B73xMo17", "Mo17")
PolarCoordHeterPlot(x=x, gt=gt, plot_lab="graph", thr=0.01, rev_log=exp(1))

coord <- PolarCoordHeterPlot(x=x, gt=gt, thr=0.01, rev_log=exp(1))
points(x=coord$x[3], coord$y[3], pch=22, cex=4, col=2)
# using random data
gt <- c("P1", "P1xP2", "P2")
set.seed(0)
x <- matrix(rnorm(150), nrow = 10, dimnames = list(paste0("M", 1:10), sample(rep(gt, 5))))
x[1:4, 1:6]
PolarCoordHeterPlot(x=x, gt=gt)
# using text style labels for the sections
PolarCoordHeterPlot(x=x, gt=gt, plot_lab="text", exp_fac=0.75)
# reverting the order of parental genotypes
PolarCoordHeterPlot(x=x, gt=c("P2", "P1xP2", "P1"), plot_lab="text", exp_fac=0.75)
# using graph style labels for the sections
PolarCoordHeterPlot(x=x, gt=c("P2", "P1xP2", "P1"), plot_lab="graph")
# coloring data points
PolarCoordHeterPlot(x=x, gt=gt, col=1:10)
# applying ANOVA P value threshold to input rows
PolarCoordHeterPlot(x=x, gt=gt, col=1:10, thr=0.5)
PolarCoordHeterPlot(x=x, gt=gt, plot_lab="graph", col=1:10, thr=0.5)
```

raw

Metabolomics data set

Description

This data set contains a matrix of log10-transformed ion intensities from a maize root metabolomics study for in total 112 metabolites in 120 samples.

Usage

raw

Format

An object of class *matrix* (inherits from *array*) with 120 rows and 112 columns.

Author(s)

Jan Lisec <jan.lisec@bam.de>

Source

<<https://onlinelibrary.wiley.com/doi/10.1111/j.1365-313X.2011.04689.x>>

RemoveFactorsByANOVA *RemoveFactorsByANOVA.*

Description

RemoveFactorsByANOVA will remove variance from data using an ANOVA model.

Usage

```
RemoveFactorsByANOVA(  
  y = NULL,  
  sam = NULL,  
  fmod = NULL,  
  kmod = NULL,  
  output = c("y_norm", "y_lm", "anova_y", "anova_y_norm", "boxplot")[1],  
  remove_outliers = 0  
)
```

Arguments

<code>y</code>	Data vector (or data matrix) to normalize (numeric + in same order as sam).
<code>sam</code>	data.frame containing the factors or numerical vars for ANOVA model.
<code>fmod</code>	Full model describing the experimental setting (provided as character string).
<code>kmod</code>	Reduced model describing all the biological factors to keep (provided as character string).
<code>output</code>	Should be <code>y_norm</code> in general but can be switched for testing.
<code>remove_outliers</code>	Should be a numeric integer <code>x</code> (with <code>\$x=0\$</code> : no effect; <code>\$x>=1\$</code> remove all values which have error <code>e</code> with <code>\$e > abs(mean + x * sd)\$</code>).

Details

See examples.

Value

Depends on output. Usually the normalized data vector (or matrix).

Examples

```
# set up sample information
sam <- data.frame(
  "GT" = gl(4, 10),
  "TR" = rep(gl(2, 5), 4),
  "Batch" = sample(gl(2, 20)),
  "Order" = sample(seq(-1, 1, length.out = 40))
)

# set up artificial measurement data
set.seed(1)
# specify main effects
m1 <- c(5, 6, 2, 9)[sam$GT] + c(-2, 2)[sam$TR]
m2 <- c(5, -6, 2, 4)[sam$GT] + c(-2, 2)[sam$TR]
# add run order bias and noise
m1 <- m1 + c(-3, 3)[sam$Batch] + 3 * sam$Order + rnorm(nrow(sam), sd = 0.5)
m2 <- m2 - 5 * sam$Order + rnorm(nrow(sam), sd = 0.8)
dat <- data.frame(m1, m2)

# apply function to remove variance
# full model incorporating all relevant factors defined in sample table
fmod <- "GT*TR+Batch+Order"
# reduced model: factors to be kept from full model; everything else will be removed from the data
kmod <- "GT*TR"
RemoveFactorsByANOVA(y = dat[, "m1"], sam = sam, fmod = fmod, kmod = kmod, output = "anova_y")
RemoveFactorsByANOVA(y = dat[, "m1"], sam = sam, fmod = fmod, kmod = kmod, output = "anova_y_norm")
```

ReplaceMissingValues *ReplaceMissingValues.*

Description

ReplaceMissingValues will replace missing values within a numeric matrix based on a principal component analysis.

Usage

```
ReplaceMissingValues(x, ncomp = 10, silent = FALSE)
```

Arguments

x	Numeric matrix.
ncomp	Number of components to be used.
silent	FALSE, suppress messages setting silent=TRUE.

Details

The ‘nipals’ algorithm is used to basically perform a PCA on the sparse matrix. Missing values are imputed based on the major components observed. Please check also the ‘impute.nipals’ function from mixOmics which should basically give the same functionality since the 04/2021 update.

Value

A matrix of similar dimensions as x without missing values.

Examples

```
# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam

idx <- apply(raw, 2, CheckForOutliers, group = sam$GT, n_sd = 5, method = "logical")
sum(idx) # 215 values would be classified as outlier using a five-sigma band
old_vals <- raw[idx] # keep outlier values for comparison
raw_filt <- raw
raw_filt[idx] <- NA
raw_means <- apply(raw, 2, function(x) {
  sapply(split(x, sam$GT), mean, na.rm = TRUE)[as.numeric(sam$GT)]
})[idx]
raw_repl <- ReplaceMissingValues(x = raw_filt)
new_vals <- raw_repl[idx]
par(mfrow = c(2, 1))
breaks <- seq(-0.7, 1.3, 0.05)
hist(raw_means - old_vals, breaks = breaks, main = "", xlab = "Outliers", las = 1)
hist(raw_means - new_vals, breaks = breaks, main = "", xlab = "Replaced values", las = 1)
```

RestrictedPCA

RestrictedPCA.

Description

RestrictedPCA combines an ANOVA based on 'fmod' and restricts a PCA using the ANOVA result as a filter.

Usage

```
RestrictedPCA(
  dat = NULL,
  sam = NULL,
  use.sam = NULL,
  group.col = NULL,
  text.col = NULL,
  fmod = NULL,
  sign.col = NULL,
  p.adjust.method = "none",
  P = 0.01,
  pcaMethods.scale = "pareto",
  n.metab.min = 20,
  ...
)
```

Arguments

<code>dat</code>	Metabolite matrix (samples x metabolites).
<code>sam</code>	Sample definition dataframe.
<code>use.sam</code>	Numeric index vector (or logical) to select specific samples to be included in the analysis or NULL to include all.
<code>group.col</code>	Column used for legend creation (column name from sam).
<code>text.col</code>	Column used for text annotation of data points (column name from sam).
<code>fmod</code>	ANOVA model to calculate before PCA.
<code>sign.col</code>	Which column(s) of the ANOVA result shall be used for P-value filtering (specify column names or leave on NULL to filter on all).
<code>p.adjust.method</code>	Method use to adjust P-values (e.g. none, BH or bonferroni).
<code>P</code>	P-value threshold used as a cutoff after P-value adjustment.
<code>pcaMethods.scale</code>	pcaMethods scale parameter (usually pareto for metabolite data).
<code>n.metab.min</code>	Minimum number of metabolites kept for PCA calculation (even if they exceed P).
<code>...</code>	Handed through to PlotMetabolitePCA .

Details

'fmod' should be something like 'GT*TR+Batch' to perform an ANOVA with these factors defined as columns in sam.

Value

Will generate a PCA plot (generated by [PlotMetabolitePCA](#) internally) restricted based on an ANOVA result based on [MetaboliteANOVA](#).

Examples

```
# load raw data and sample description
raw <- MetabolomicsBasics::raw
sam <- MetabolomicsBasics::sam
# standard behavior
RestrictedPCA(dat = raw, sam = sam, group.col = "GT")
## Not run:
# apply multiple testing using a strict P-value cutoff,
# dont show a legend but plot group mean values and sd's as overlay
RestrictedPCA(
  dat = raw, sam = sam, group.col = "GT", p.adjust.method = "BH", P = 10^-10,
  fmod = "GT+Batch+Order", sign.col = "GT", medsd = T, legend.x = NULL
)
# limit to a subset of samples, switching the ANOVA selection of by setting P=1
# and adding text (from \code{sam}) to each data point
RestrictedPCA(
  dat = raw, sam = sam, use.sam = which(sam$GT %in% c("Mo17", "B73")), group.col = "GT",
  fmod = "GT+Batch+Order", P = 1, sign.col = "GT", legend.x = NULL, text.col = "Batch"
)

## End(Not run)
```

sam

Sample table

Description

This data frame contains the sample definition of 120 samples according to the rows of [raw](#).

Usage

```
sam
```

Format

An object of class `data.frame` with 120 rows and 10 columns.

Author(s)

Jan Lisec <jan.lisec@bam.de>

Source

<<https://onlinelibrary.wiley.com/doi/10.1111/j.1365-313X.2011.04689.x>>

spectra_format_converter

spectra_format_converter.

Description

spectra_format_converter will generate a matrix with mz and int columns out of a text representation of a spectrum.

Usage

```
spectra_format_converter(txt = NULL, m_prec = 3, i_prec = 0)
```

Arguments

txt	Sample table.
m_prec	Mass precision of output spectrum.
i_prec	Intensity precision of output spectrum.

Details

See examples.

Value

Matrix with mz and int columns.

Examples

```
spectra_format_converter(txt = "57.1:100 58.0001:10")  
spectra_format_converter(txt = "58.0001:10 57.1:100", m_prec = 4)
```

sumformula_from_CAS	<i>sumformula_from_CAS.</i>
---------------------	-----------------------------

Description

sumformula_from_CAS will try to get a chemical sum formula from a CAS ID.

Usage

```
sumformula_from_CAS(x = NULL)
```

Arguments

x	Vector of CAS IDs.
---	--------------------

Details

tbd.

Value

A character vector of length input vector.

Examples

```
## Not run:  
x <- readLines("C:/Users/jlisec/Documents/Francesco Russo/RECTOX/RECTOX_GC-EI-MS_CASRN")  
sf <- sumformula_from_CAS(x = x)  
  
## End(Not run)
```

unique_labels	<i>unique_labels.</i>
---------------	-----------------------

Description

unique_labels will generate a dataframe with color and plotting character specification out of a sample table definition.

Usage

```
unique_labels(sam = NULL, g = NULL)
```

Arguments

sam	Sample table.
g	Either column name from sam containing factor column or factor of same length as sam.

Details

If a color/symbol specification exists for a sample set containing replicate groups this function will help in retrieving this information per group which is useful in boxplot or legend functions (cf. examples).

Value

Dataframe with group levels names and their color and plotting character specification.

Examples

```
sam <- MetabolomicsBasics::sam  
unique_labels(sam = sam, g = "GT")
```

unique_subformula_masses
unique_subformula_masses

Description

unique_subformula_masses will generate a numeric vector of potential sub formula masses regarding a chemical formula as input.

Usage

```
unique_subformula_masses(fml, names = TRUE, check_validity = FALSE)
```

Arguments

fml	Chemical formula.
names	Return respective sub formulas as vector names.
check_validity	Filter for chemically valid formulas.

Details

In mass spectrometry precursor masses are often fragmented and these fragments are recorded as MS² spectra. A frequent task is then to compute potential chemical formulas for the obtained MS² masses. The function unique_subformula_masses follows the reverse approach. It allows to calculate all masses that could be potential breakdown products of a precursor formula

Value

A named numeric vector. The names are the sub formulas for the calculated exact masses given as numeric.

Examples

```
# specify a formula and calculate all potential combinatorial masses
fml <- c("C6H12O6", "C11H16N04PS", "C24H51O4P")[1]
tmp <- unique_subformula_masses(fml = fml)
length(tmp); any(duplicated(tmp))
hist(tmp, breaks=seq(floor(min(tmp))-1, ceiling(max(tmp))), main=fml)
# do the same as above but check for chemical plausibility
tmp2 <- unique_subformula_masses(fml = fml, check_validity=TRUE)
length(tmp2)
hist(tmp2, breaks=seq(floor(min(tmp2))-1, ceiling(max(tmp2))), main=fml)
mz <- 147
tmp[abs(tmp-mz)<0.5]
tmp2[abs(tmp2-mz)<0.5]
```

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