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ChemoSpecUtils-package .................................................. 2
check4Gaps ................................................................. 2
checkForPackageWithVersion .......................................... 4
cchkSpectra ............................................................... 4
Functions Supporting Packages ChemoSpec and ChemoSpec2D

Description

Functions supporting the packages ChemoSpec and ChemoSpec2D.

Author(s)

Bryan A. Hanson.
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check4Gaps

Check for Discontinuities (Gaps) in a Vector & Optionally Make a Plot

Description

The basic procedure is to compare \( x[n + 1] - x[n] \) for successive values of \( n \). When this value jumps, there is a gap which is flagged. \( \text{beg.indx} \) and \( \text{end.indx} \) will always be contiguous as indices must be; it is the \( x \) values that jump or have the gap. The indices are provided as they are more convenient in some programming contexts. If not assigned, the result appears at the console.

Usage

\[
\text{check4Gaps}(x, y = \text{NULL}, \text{silent} = \text{FALSE}, \text{tol} = \text{NULL}, \ldots)
\]
check4Gaps

Arguments

x  A numeric vector to be checked for gaps.
y  An optional vector of y-values which correspond to the x values. Only used in ChemoSpec. If provided, a plot will be made in the style of a Spectra object showing the gap(s).
silent  Logical indicating a "no gap" message should not be reported to the console. Important because check4Gaps is called iteratively by other functions.
tol  A number indicating the tolerance for checking to see if the step between successive x values are the same. Depending upon how the x values are stored and rounded, you may need to change the value of tol to avoid flagging trivial "gaps". If NULL, a value is chosen which is just above the median difference between x values.

...  Other parameters to be passed to the plot routines if y is provided, e.g. xlim.

Value

A data frame giving the data chunks found, with one chunk per row. Also a plot if y is provided. In the event there are no gaps found, a data frame with one row is returned. The data frame has columns as follows:

beg.freq  The first frequency value in a given data chunk.
end.freq  The last frequency value in a given data chunk.
size  The length (in frequency units) of the data chunk.
beg.indx  The index of the first frequency value in the data chunk.
end.indx  The index of the last frequency value in the data chunk.

Author(s)

Bryan A. Hanson, DePauw University.

See Also

sumSpectra which make extensive use of this function.

Examples

x <- seq(0, 2 * pi, 0.1)
y <- sin(x)
remove <- c(8:11, 40:45)
x <- x[-remove]
y <- y[-remove]
gaps <- check4Gaps(x, tol = 0.11) # tol just larger than orig spacing
gaps
gaps <- check4Gaps(x, y, tol = 0.11) # show a plot if y given
checkForPackageWithVersion

*Check for an Installed Package with a Particular Version or Newer*

**Description**

Utility function for making sure a package is available with a particular version or newer.

**Usage**

```
checkForPackageWithVersion(pkg, vers)
```

**Arguments**

- `pkg` Character. The name of the package to check.
- `vers` Character. The minimum acceptable version of the package. Will only be checked to the major.minor level.

**Value**

If successful, `TRUE` is return invisibly. Stops if there is a problem.

chkSpectra

*Verify the Integrity of a Spectra or Spectra2D Object*

**Description**

Utility function to verify that the structure of a Spectra or Spectra2D object is internally consistent. This function should be used after manual editing of these objects. However, in most cases rather than directly editing these objects, one should modify them via:

- `removeFreq`
- `removeSample`
- `removeGroup`

**Usage**

```
chkSpectra(spectra, confirm = FALSE)
```

**Arguments**

- `spectra` An object of S3 class Spectra or Spectra2D.
- `confirm` Logical indicating whether or not to write the results to the console, as would be desirable for interactive use.
**colorSymbol**

**Value**

None. When used at the console, and the object is OK, no message is written unless `confirm = TRUE`. At the console, if there is a problem, messages are issued regardless of the value of `confirm`.

**Author(s)**

Bryan A. Hanson, DePauw University.

**Examples**

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)
  chkSpectra(SrE.IR)
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
  chkSpectra(MUD1)
}
```

**Description**

In ChemoSpec and ChemoSpec2D, the user may use any color name/format known to R. The current color scheme of a Spectra or Spectra2D object may be determined using `sumGroups` or `sumSpectra`. The colors can also be queried and changed using `conColScheme`.

**Format**

Colors are stored as character vectors and symbols as numeric vectors.

**Details**

An important fact to keep in mind is that most people with normal vision cannot distinguish more than about 8-12 colors, and doing so depends upon the viewing circumstances: if on paper, printer, ink and paper type all matter, and if on a screen, the background color makes a big difference. Further, color-blind individuals have additional challenges. A great discussion of color issues can be found in the colorspace package. The Polychrome package has further discussion and utilities for choosing qualitative colorschemes, including those for color-blind individuals.

ChemoSpec, but not ChemoSpec2D, can also create plots using the built-in symbols and lower case letters. This is useful for color-blind individuals, plots in rgl which can’t plot regular symbols, and plots for where there are more groups than could be reasonably coded in color. A good discussion of which symbols are most readily distinguished can be found in Robinson: "Good Plot Symbols by Default" Journal of Computational and Graphical Statistics DOI: 10.1080/10618600.2019.1637746
ChemoSpecUtils supplies four color/symbol schemes for your consideration. If the particular order of colors in any of these does not suit your needs, you can always choose the ones you want, and/or rearrange the order, or simply provide your own.

- The colors and symbols produced by `gr.cols = "auto"` in the import functions.
- `Col8` provides eight unique colors. These are more saturated than the automatic colors.
- `Col12` provides a mostly paired set of 12 unique colors suitable for groups that come in pairs.
- `Col7` provides seven color-blind friendly colors. These can be visualized at https://projects.susielu.com/viz-palette by using the hex codes obtained by typing `data(Col7); Col7` in the R console.

**Author(s)**

Bryan A. Hanson, DePauw University.

**Examples**

```r
# Make a plot showing all the built-in color options

data(Col7)
data(Col12)
data(Sym12)
data(Col8)
data(Sym8)
auto <- RColorBrewer::brewer.pal(8, "Set1")

sp <- 0.75 # space between major plot elements
tsp <- 0.15 # additional space between points and color swatches/descriptive text
h <- 0.25 # height of the swatch
y <- 0.0 # bottom of the plot, the reference point

# empty plot
plot(1:12, rep(0.0, 12),
     type = "n", yaxt = "n", xaxt = "n", bty = "n",
     xlab = "", ylab = "", ylim = c(0, 3.5))

text(6.5, y + h + tsp * 4 + sp * 3.5,
     labels = "Automatic Color & Symbol Options", cex = 1.25, font = 2)

# Col12
for (i in 1:12) {
  rect(i - 0.5, y, i + 0.5, y + h, border = NA, col = Col12[i])
}
points(1:12, rep(y + h + tsp, 12), pch = Sym12)

# Col8
for (i in 1:8) {
```

```
conColScheme

Change the Color Scheme of a Spectra or Spectra2D Object

Description

This function permits you to change the color scheme of an existing Spectra or Spectra2D object.

Usage

conColScheme(spectra, new.cols = NULL, silent = FALSE)

Arguments

spectra An object of S3 class Spectra or Spectra2D.
new.cols A character vector giving the new color values, of length(unique(spectra$colors)). If not provided, the function will print the old values for reference.
silent Logical. If TRUE, suppresses all reporting.

Value

spectra An updated object of S3 class Spectra or Spectra2D.

See Also

For a discussion of general issues of color, see colorSymbol.
Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(metMUD1)

  sumSpectra(metMUD1)
  newSpec <- conColScheme(metMUD1) # reports old colors
  newSpec <- conColScheme(metMUD1, new = c("pink", "violet"))
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)

  sumSpectra(MUD1)
  newSpec <- conColScheme(MUD1) # reports old colors
  newSpec <- conColScheme(MUD1, new = c("pink", "violet"))
}
```

**hcaScores**

_HCA on PCA/MIA/PARAFAC scores from a Spectra or Spectra2D Object_

**Description**

A wrapper which performs HCA on the scores from a PCA of a Spectra object or POP/MIA/PARAFAC of a Spectra2D object. Many methods for computing the clusters and distances are available.

**Usage**

```r
hcaScores(
  spectra, 
  so, 
  scores = c(1:5), 
  c.method = "complete", 
  d.method = "euclidean", 
  use.sym = FALSE, 
  leg.loc = "topright", 
  ...
)
```

**Arguments**

- `spectra` An object of S3 class Spectra or Spectra2D object.
- `so` "Score Object" One of the following:
  - An object of class prcomp, created by ChemoSpec functions c_pcaSpectra, r_pcaSpectra, irlba_pcaSpectra or s_pcaSpectra.
hcaScores

- An object of class mia produced by function miaSpectra2D.
- An object of class parafac produced by function pfacSpectra2D.
- An object of class pop produced by function popSpectra2D.

Any of the above score objects will have been modified to include a list element called $method, a character string describing the pre-processing carried out and the type of PCA performed (used to annotate the plot).

scores A vector of integers specifying the components (scores) to plot.
c.method A character string describing the clustering method; must be acceptable to hclust.
d.method A character string describing the distance calculation method; must be acceptable as a method in rowDist.
use.sym A logical; if true, use no color and use lower-case letters to indicate group membership. Applies only to Spectra objects.
leg.loc Character; if “none” no legend will be drawn. Otherwise, any string acceptable to legend.
...
Additional parameters to be passed to the plotting functions.

Value

A list, containing an object of class hclust and an object of class dendrogram. The side effect is a plot.

Author(s)

Bryan A. Hanson, DePauw University.

See Also

hclust for the underlying function. See hcaSpectra for HCA of the entire data set stored in the Spectra object.

Examples

if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
data(metMUD1)

  pca <- c_pcaSpectra(metMUD1)
hca <- hcaScores(metMUD1, pca, main = "metMUD1 NMR Data PCA Scores")
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
data(MUD1)

  mia <- miaSpectra2D(MUD1)
hca <- hcaScores(MUD1, mia, scores = 1:2, main = "MUD1 MIA Scores")
  set.seed(123)
pfac <- pfacSpectra2D(MUD1, parallel = FALSE, nfac = 2)
hca <- hcaScores(MUD1, pfac, scores = 1:2, main = "MUD1 PARAFAC Scores")

plotScores

Plot Scores from PCA, MIA or PARAFAC Analysis of a Spectra or Spectra2D Object

Description

Plots the requested scores using the color scheme derived from the Spectra or Spectra2D object. Options are provided to add confidence ellipses for each group in the object. The ellipses may be robust or classical. Option to label the extreme points provided.

Usage

plotScores(
spectra,  
so,     
pcs = c(1, 2),     
ellipse = "none",  
tol = "none",     
use.sym = FALSE,  
leg.loc = "topright",  
...  
)

Arguments

spectra An object of S3 class Spectra or Spectra2D object.
so "Score Object" One of the following:

- An object of class prcomp, created by ChemoSpec functions c_pcaSpectra, r_pcaSpectra, irlba_pcaSpectra or s_pcaSpectra.
- An object of class mia produced by function miaSpectra2D.
- An object of class parafac produced by function pfacSpectra2D.
- An object of class pop produced by function popSpectra2D.

Any of the above score objects will have been modified to include a list element called $method, a character string describing the pre-processing carried out and the type of PCA performed (used to annotate the plot).

pcs A vector of two integers specifying the components (scores) to plot.
ellipse A character vector specifying the type of ellipses to be plotted. One of c("both", "none", "cls", "rob"). cls specifies classical confidence ellipses, rob specifies robust confidence ellipses. An ellipse is drawn for each group unless there are three or fewer samples in the group.
tol A number describing the fraction of points to be labeled. \( tol = 1.0 \) labels all the points; \( tol = 0.05 \) labels the most extreme 5 percent. Set to "none" to completely suppress labels.

use.sym A logical; if TRUE, the color scheme is set to black and the points plotted with symbols. Applies only to \texttt{Spectra} objects.

leg.loc Character; if "none" no legend will be drawn. Otherwise, any string acceptable to \texttt{legend}.

additional parameters to be passed to the plotting functions.

Value

None. Side effect is a plot.

Author(s)

Bryan A. Hanson, DePauw University.

Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(metMUD1)

  pca <- c_pcaSpectra(metMUD1)
  plotScores(metMUD1, pca,
    main = "metMUD1 NMR Data",
    pcs = c(1, 2), ellipse = "cls", tol = 0.05
  )
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)

  res <- miaSpectra2D(MUD1)
  plotScores(MUD1, res, main = "MIA Scores", tol = 0.1, ellipse = "cls")

  set.seed(123)
  res <- pfacSpectra2D(MUD1, parallel = FALSE, nfac = 2)
  plotScores(MUD1, res, tol = 0.1, leg.loc = "bottomright", main = "PARAFAC Score Plot")
}
```
plotScree

Description

Functions that draw a traditional scree plot, or an alternative style that is perhaps more informative. These plots illustrate the variance explained by each component in a PCA or MIA analysis.

Usage

plotScree(pca, style = "alt", ...)

Arguments

- pca
  - Either: an object of class prcomp, modified to include a list element called $method, a character string describing the pre-processing carried out and the type of PCA performed (it appears on the plot). This is automatically provided if ChemoSpec functions c_pcaSpectra or r_pcaSpectra were used to create pca.
  - An object of class mia produced by function miaSpectra2D.
- style
  - Character. One of c("trad","alt") giving the style of plot desired (traditional or alternative). "trad" is not supported for mia objects.
- ...
  - Additional parameters to be passed to plotting functions.

Value

None. Side effect is a plot.

Author(s)

Bryan A. Hanson, DePauw University.

References

The idea for the alternative style plot came from the NIR-Quimiometria blog by jrcuesta, at [https://nir-quimiometria.blogspot.com/2012/02/pca-for-nir-spectrapart-004-projections.html](https://nir-quimiometria.blogspot.com/2012/02/pca-for-nir-spectrapart-004-projections.html)

Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(metMUD1)

  pca <- c_pcaSpectra(metMUD1)
  plotScree(pca, style = "trad")
  plotScree(pca, style = "alt")
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
```
```
mia <- miaSpectra2D(MUD1)
plotScree(mia, style = "alt")
```

---

**removeFreq**

*Remove Frequencies from a Spectra or Spectra2D Object*

**Description**

This function removes specified frequencies from a **Spectra** or **Spectra2D** object. For instance, one might want to remove regions lacking any useful information (to reduce the data size), remove regions with large interfering peaks (e.g. the water peak in 1H NMR) or simply focus on a region of interest.

**Usage**

```r
removeFreq(spectra, rem.freq = NULL, remF2 = NULL, remF1 = NULL)
```

**Arguments**

- **spectra**: An object of S3 class **Spectra** or **Spectra2D** from which to remove frequencies.
- **rem.freq**: For a **Spectra** object, a vector of logicals. **rem.freq** can be any valid R statement that leads to a vector of logicals (must be of length(Spectra$freq)). This vector should be TRUE for frequencies you want to be removed and FALSE for those frequencies which will be kept. In the examples, the | and & operators may seem backward in a sense, but R evaluates them one at a time and then combines them to give the desired result. You may wish to look at **Comparison** and **Logic**. See the examples. *In addition, since January 2020 rem.freq may be a formula as described below.*
- **remF2**: Applies to **Spectra2D** objects. A formula giving the range of frequencies to be extracted. May include "low" or "high" representing the extremes of the spectra. Values outside the range of F2 are tolerated without notice and are handled as min or max. See the examples.
- **remF1**: As for remF2.

**Value**

An object of S3 class **Spectra** or **Spectra2D**.

**Modifying Spectra2D Objects**

Regarding **Spectra2D** objects, one cannot remove frequencies from the interior of a 2D NMR data set and expect to get a meaningful contour plot, because doing so puts unrelated peaks adjacent in the data set. This would lead to contours being drawn that don’t exist in the original data set. However, one can remove data from the interior and run a PARAFAC analysis on the result, using the spectrum as an abstract object (that is, the spectrum may not plottable, but the resulting scores are still meaningful).
Author(s)

Bryan A. Hanson, DePauw University.

See Also

removePeaks2D for another way to remove data.

Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)
  sumSpectra(SrE.IR)

  # Examples where rem.freq is a logical vector
  # Remove frequencies from one end:
  newIR <- removeFreq(SrE.IR, rem.freq = SrE.IR$freq > 3500)

  # Remove frequencies from both ends at once:
  newIR <- removeFreq(SrE.IR, rem.freq = SrE.IR$freq > 3500 |
                      SrE.IR$freq < 800)

  # Remove frequencies from the middle:
  newIR <- removeFreq(SrE.IR, rem.freq = SrE.IR$freq > 800 &
                      SrE.IR$freq < 1000)

  # The logic of this last one is as follows. Any values
  # that are TRUE will be removed.
  values <- 1:7
  values > 2
  values < 6
  values > 2 & values < 6

  # Examples where rem.freq is a formula
  # Remove frequencies from one end:
  newIR <- removeFreq(SrE.IR, rem.freq = 3500 ~ high)

  # Remove frequencies from both ends is a two step process with formulas:
  newIR <- removeFreq(SrE.IR, rem.freq = 3500 ~ high)
  newIR <- removeFreq(newIR, rem.freq = low ~ 800)

  # Remove frequencies from the middle:
  newIR <- removeFreq(SrE.IR, rem.freq = 800 ~ 1000)

  # After any of these, inspect the results:
  sumSpectra(newIR)
}
```

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
```
data(MUD1)

plotSpectra2D(MUD1,
    which = 7, lvls = seq(-1, 1, by = 0.2),
    main = "MUD1 Sample 7: Complete Data Set"
)

MUD1a <- removeFreq(MUD1, remF2 = 2 ~ 4)
sumSpectra(MUD1a) # cannot plot, results would be misleading

MUD1b <- removeFreq(MUD1, remF1 = low ~ 20)
sumSpectra(MUD1b)
plotSpectra2D(MUD1b,
    which = 7, lvls = seq(-1, 1, by = 0.2),
    main = "MUD1 Sample 7\nRemoved Frequencies: F1 low ~ 20"
)

MUD1c <- removeFreq(MUD1, remF2 = low ~ 2)
sumSpectra(MUD1c)
plotSpectra2D(MUD1c,
    which = 7, lvls = seq(-1, 1, by = 0.2),
    main = "MUD1 Sample 7\nRemoved Frequencies: F2 low ~ 2"
)

MUD1d <- removeFreq(MUD1, remF2 = 3 ~ high, remF1 = 45 ~ 55)
sumSpectra(MUD1d) # not plotted, results would be misleading

---

**removeGroup**

**Remove a Group from a Spectra or Spectra2D Object**

**Description**

Removes specified groups from a **Spectra** or **Spectra2D** object.

**Usage**

```r
removeGroup(spectra, rem.group)
```

**Arguments**

- **spectra** An object of S3 class **Spectra** or **Spectra2D**.
- **rem.group** A character vector (handled as a regex) giving the groups to be removed.

**Details**

This function will report if extra data elements are found. These will probably need to be edited manually. The indices reported to the console can be helpful in this regard.
removeSample

If `rem.group` is a character vector, the sample names are grepped for the corresponding values. Remember that the grepping process is greedy, i.e. grepping for "XY" find not only "XY" but also "XYZ".

Unused levels in $groups$ are dropped.

Value

An object of S3 class `Spectra` or `Spectra2D`.

Author(s)

Bryan A. Hanson, DePauw University.

Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)

  sumGroups(SrE.IR)
  SrE.IRa <- removeGroup(SrE.IR, rem.group = "pSrE")
  sumGroups(SrE.IRa)
}
if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)

  sumGroups(MUD1)
  MUD1a <- removeGroup(MUD1, rem.group = "Ether")
  sumGroups(MUD1a)
}
```

removeSample Remove Samples from a Spectra or Spectra2D Object

Description

Removes specified samples from a `Spectra` or `Spectra2D` object.

Usage

```r
removeSample(spectra, rem.sam)
```

Arguments

- `spectra` An object of S3 class `Spectra` or `Spectra2D`.
- `rem.sam` Either an integer vector specifying the samples to be removed, or a character vector (handled as a regex) giving the sample names to be removed.
removeSample

Details

This function will report if extra data elements are found. These will probably need to be edited manually. The indices reported to the console can be helpful in this regard.

If rem.sam is a character vector, the sample names are grepped for the corresponding values. Remember that the grepping process is greedy, i.e. grepping for "XY" find not only "XY" but also "XYZ".

Value

An object of S3 class Spectra or Spectra2D.

Author(s)

Bryan A. Hanson, DePauw University.

Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)
  
  # Remove the 9th spectrum/sample:
  SrE.IR$names
  SrE.IRa <- removeSample(SrE.IR, rem.sam = 9)
  SrE.IRa$names
  
  # Removes a spectrum/sample with this exact name:
  SrE.IRb <- removeSample(SrE.IR, rem.sam = "NW_adSrE")
  SrE.IRb$names
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
  
  # Removes the 5th spectrum:
  MUD1$names
  MUD1a <- removeSample(MUD1, rem.sam = 5)
  MUD1a$names
  
  # Removes a spectrum/sample with this exact name:
  MUD1$names
  MUD1b <- removeSample(MUD1, rem.sam = "Ether_3")
  MUD1b$names
}
```
Description

This function computes the distance between rows of a matrix using a number of methods. It is primarily a wrapper for `Dist` which provides many options. However, cosine distance is calculated locally. See the reference for an excellent summary of distances and similarities. Keep in mind that distances are always positive by definition. Further, in the literature one can find the same distance defined different ways. For instance, the definition of the "pearson" and "correlation" distances differs slightly between the reference below and `Dist`. So please study the definitions carefully to get the one you want. The example illustrates the behavior of some common distance definitions. Notice that "pearson" and "cosine" are mathemtically identical for the particular definition of "pearson" used by `Dist`.

Usage

```r
rowDist(x, method)
```

Arguments

- `x`: A matrix whose rows will be used for the distance calculation.
- `method`: Character; one of "cosine", "euclidean", "maximum", "manhattan", "canberra", "binary", "pearson", "correlation", "spearman", "kendall", "abspearson", "abscorrelation".

Value

An object of class `dist`.

Author(s)

Bryan A. Hanson, DePauw University.

References


Examples

```r
# This examples imagines spectra as a series of vectors
# on a half unit circle.
# 1. Compute half of a unit circle
theta <- seq(0, pi, length = 100)
x = cos(theta)
y = sin(theta)
```
# 2. Compute some illustrative vectors
# Get tail/origin & tip/head coordinates
lt <- length(theta)
set.seed(6)
tips <- theta[c(1, sample(2:100, 5))]
x0 <- y0 <- rep(0.0, lt) # tail/origin at 0,0
x1 <- cos(tips) # tips/heads
y1 <- sin(tips)

# 3. Compute the distance functions
# Bounded distances
RDcor <- rep(NA_real_, lt) # correlation distance
RDpea <- rep(NA_real_, lt) # pearson distance
RDabp <- rep(NA_real_, lt) # abspearson distance
RDcos <- rep(NA_real_, lt) # cosine distance

# Unbounded distances
RDeuc <- rep(NA_real_, lt) # Euclidean distance
RDman <- rep(NA_real_, lt) # manhattan distance

# Compute all
np <- 5
refVec <- c(seq(0.0, x[1], length.out = np), seq(0.0, y[1], length.out = np))
for (i in 1:lt) {
  Vec <- c(seq(0.0, x[i], length.out = np), seq(0.0, y[i], length.out = np))
  M <- matrix(c(refVec, Vec), nrow = 2, byrow = TRUE)
  RDman[i] <- rowDist(M, method = "manhattan")
  RDeuc[i] <- rowDist(M, method = "euclidean")
  RDcos[i] <- rowDist(M, method = "cosine")
  RDcor[i] <- rowDist(M, method = "correlation")
  RDpea[i] <- rowDist(M, method = "pearson")
  RDabp[i] <- rowDist(M, method = "abspearson")
}

# 4. Plots
# a. Unit circle w/representative vectors/spectra
plot.new()
plot.window(xlim = c(-1, 1), ylim = c(0, 1), asp = 1)
title(main = "Representative 'Spectral' Vectors on a Unit Half Circle\nReference Vector in Red",

sub = "Each 'spectrum' is represented by a series of x, y points")
lines(x, y, col = "gray") # draw half circle
lines(x = x[c(1, 100)], y = y[c(1, 100)], col = "gray") # line across bottom
arrows(x0, y0, x1, y1, angle = 5) # add arrows & a red reference vector
arrows(x0[1], y0[1], x1[1], y1[1], col = "red", angle = 5, lwd = 2)

# b. Distances
degrees <- theta*180/pi
plot(degrees, RDman, type = "l",
xlab = "Angle Between Spectral Vectors and Reference Vector in Degrees",
ylab = "Distance",
main = "Spectral Distance Comparisons\nUsing ChemoSpecUtils::rowDist")
abline(h = c(1.0, 2.0), col = "gray")

# 19
lines(degrees, RDeuc, col = "blue")
lines(degrees, RDcos, col = "green", lwd = 4)
lines(degrees, RDcor, col = "red")
lines(degrees, RDabp, col = "black", lty = 2)
lines(degrees, RDpea, col = "black", lty = 3)

leg.txt <- c("manhattan", "euclidean", "correlation", "cosine", "pearson", "abspearson")
leg.col <- c("black", "blue", "red", "green", "black", "black")
leg.lwd <- c(1, 1, 1, 1, 4, 1)
leg.lty <- c(1, 1, 1, 1, 3, 2)
legend("topleft", legend = leg.txt, col = leg.col, lwd = leg.lwd, lty = leg.lty)

---

**sampleDist**

Compute the Distances Between Samples in a Spectra or Spectra2D Object

**Description**

Compute the distances between samples in a Spectra or Spectra2D object. This is a means to quantify the similarity between samples. A heat map style plot is an option.

**Usage**

```r
sampleDist(spectra, method = "pearson", plot = TRUE, ...)
```

**Arguments**

- `spectra`: An object of S3 class Spectra or Spectra2D.
- `method`: Character. A string giving the distance method. See `rowDist` for options.
- `plot`: Logical. Shall a level plot (heat map) be made?
- `...`: Arguments to be passed to the plotting function.

**Value**

A numeric matrix giving the distances between the samples.

**Author(s)**

Bryan A. Hanson, DePauw University.

**See Also**

For Spectra objects, see `plotSpectraDist` which compares all spectra to a single reference spectrum.
Examples

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
    library("ChemoSpec")
    library("lattice")
    data(SrE.IR)
    SrE.dmatrix <- sampleDist(SrE.IR, # cosine distance bounded on [0...2]
        method = "cosine",
        main = "SrE.IR Cosine Distance Between Samples"
    )
    SrE.dmatrix <- sampleDist(SrE.IR, # abspearson distance bounded on [0...1]
        method = "abspearson",
        main = "SrE.IR Absolute Pearson Distance Between Samples"
    )
    SrE.dmatrix <- sampleDist(SrE.IR, # euclidean distance unbounded
        method = "euclidean",
        main = "SrE.IR Euclidean Distance Between Samples"
    )
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
    library("ChemoSpec2D")
    library("lattice")
    data(MUD1)
    MUD1.dmatrix <- sampleDist(MUD1,
        method = "cosine",
        main = "MUD1 Cosine Distance Between Samples"
    )
}
```

sumGroups

---

`sumGroups` **Summarize the Group Membership of a Spectra or Spectra2D Object**

Description

This function summarizes the group membership of a Spectra or Spectra2D object.

Usage

`sumGroups(spectra)`

Arguments

- `spectra` An object of S3 class Spectra or Spectra2D whose group membership information is desired.
sumSpectra

Value
A data frame as follows. Note that if there are groups with no members these are dropped.

- **group**: The name of the group.
- **no.**: The number in the group.
- **color**: The color assigned to the group.
- **symbol**: The symbol assigned to the group. Spectra objects only.
- **alt.symbol**: The alternative symbol assigned to the group. Spectra objects only.

Author(s)
Bryan A. Hanson, DePauw University.

See Also
To summarize the entire object, sumSpectra.

Examples
```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)
  sumGroups(SrE.IR)
}
if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
  sumGroups(MUD1)
}
```

sumSpectra

Summarize a Spectra or Spectra2D Object

Description
Provides a summary of a Spectra or Spectra2D object, essentially a more spectroscopist-friendly version of `str()`.

Usage
```
sumSpectra(spectra, ...)
```

Arguments
- **spectra**: An object of S3 class Spectra or Spectra2D whose group membership information is desired.
- **...**: Arguments to be passed downstream. Main use is to pass a value for tol to function check4Gaps when using ChemoSpec. Not used in ChemoSpec2D.
**updateGroups**

**Details**

Prior to summarizing, `chkSpectra` is run with `confirm = FALSE`. If there are problems, warnings are issued to the console and the summary is not done. The `Spectra` or `Spectra2D` object is checked to see if it contains data elements beyond what is required. If so, these extra elements are reported to the console.

**Value**

None. Results printed at console.

**Author(s)**

Bryan A. Hanson, DePauw University.

**Examples**

```r
if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(SrE.IR)
  sumSpectra(SrE.IR)
}
if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
  sumSpectra(MUD1)
}
```

---

**updateGroups**

*Update Group Names in a Spectra or Spectra2D Object*

**Description**

A convenience function that can be used to update (change) group names. The default group names come from the `gr.crit` argument in the import functions `files2SpectraObject`, `matrix2SpectraObject` or `files2Spectra2DObject`. In some cases `gr.crit` may have complex regex patterns, and this function makes updating them to more appropriate/more readable strings easier.

**Usage**

```r
updateGroups(spectra, new.grps = NULL, silent = FALSE)
```

**Arguments**

- `spectra` An object of S3 class `Spectra` or `Spectra2D`.
- `new.grps` A vector of character values giving the new group names. The new values must correspond to the order of the old values. This vector should give the unique values only (so, it should have `length(unique(spectra$groups))`). If not provided, the function will print the old values for reference.
updateGroups

silent Logical. If TRUE, suppresses all reporting.

Value

spectra An updated object of S3 class Spectra or Spectra2D.

Examples

if (checkForPackageWithVersion("ChemoSpec", "5.1")) {
  library("ChemoSpec")
  data(metMUD1)
  metMUD1a <- updateGroups(metMUD1) # reports old groups
  metMUD1a <- updateGroups(metMUD1, new.grps = c("C", "T"))
}

if (checkForPackageWithVersion("ChemoSpec2D", "0.3")) {
  library("ChemoSpec2D")
  data(MUD1)
  MUD1a <- updateGroups(MUD1, new.grps = c("control", "treatment"))
}
## Index

<table>
<thead>
<tr>
<th>* classes</th>
<th>Col8 (colorSymbol), 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>chkSpectra, 4</td>
<td>ColorScheme (colorSymbol), 5</td>
</tr>
<tr>
<td>* cluster</td>
<td>colorSymbol, 5</td>
</tr>
<tr>
<td>hcaScores, 8</td>
<td>conColScheme, 5, 7</td>
</tr>
<tr>
<td>* color</td>
<td>dendrogram, 9</td>
</tr>
<tr>
<td>colorSymbol, 5</td>
<td>Dist, 18</td>
</tr>
<tr>
<td>conColScheme, 7</td>
<td>files2Spectra2DObject, 23</td>
</tr>
<tr>
<td>* datasets</td>
<td>files2SpectraObject, 23</td>
</tr>
<tr>
<td>colorSymbol, 5</td>
<td>hcaScores, 8</td>
</tr>
<tr>
<td>* hplot</td>
<td>hcaSpectra, 9</td>
</tr>
<tr>
<td>plotScores, 10</td>
<td>hclust, 9</td>
</tr>
<tr>
<td>plotScree, 11</td>
<td>irlba_pcaSpectra, 8, 10</td>
</tr>
<tr>
<td>sampleDist, 20</td>
<td>legend, 9, 11</td>
</tr>
<tr>
<td>* multivariate</td>
<td>Logic, 13</td>
</tr>
<tr>
<td>hcaScores, 8</td>
<td>matrix2SpectraObject, 23</td>
</tr>
<tr>
<td>plotScores, 10</td>
<td>miaSpectra2D, 9, 10</td>
</tr>
<tr>
<td>plotScree, 11</td>
<td>pfacSpectra2D, 9, 10</td>
</tr>
<tr>
<td>* robust</td>
<td>plotScores, 10</td>
</tr>
<tr>
<td>plotScores, 10</td>
<td>plotScree, 11</td>
</tr>
<tr>
<td>* utilities</td>
<td>plotSpectraDist, 20</td>
</tr>
<tr>
<td>check4Gaps, 2</td>
<td>popSpectra2D, 9, 10</td>
</tr>
<tr>
<td>chkSpectra, 4</td>
<td>prcomp, 8, 10, 12</td>
</tr>
<tr>
<td>colorSymbol, 5</td>
<td>removeFreq, 4, 13</td>
</tr>
<tr>
<td>conColScheme, 7</td>
<td>removeGroup, 4, 15</td>
</tr>
<tr>
<td>removeFreq, 13</td>
<td>removePeaks2D, 14</td>
</tr>
<tr>
<td>removeGroup, 15</td>
<td>removeSample, 4, 16</td>
</tr>
<tr>
<td>removeSample, 16</td>
<td>rowDist, 9, 18, 20</td>
</tr>
<tr>
<td>rowDist, 18</td>
<td>s_pcaSpectra, 8, 10</td>
</tr>
<tr>
<td>sumGroups, 21</td>
<td>sampleDist, 20</td>
</tr>
<tr>
<td>sumSpectra, 22</td>
<td>Spectra, 3–5, 7–11, 13, 15–17, 20–24</td>
</tr>
</tbody>
</table>
Spectra2D, 4, 5, 7, 8, 10, 13, 15–17, 20–24
sumGroups, 5, 21
sumSpectra, 3, 5, 22, 22
Sym12 (colorSymbol), 5
Sym8 (colorSymbol), 5

updateGroups, 23