

Package ‘hyperSpec’

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Type Package

Title Work with Hyperspectral Data, i.e. Spectra + Meta Information
(Spatial, Time, Concentration, ...)

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Description Comfortable ways to work with hyperspectral data sets.

I.e. spatially or time-resolved spectra, or spectra with any other kind of information associated with each of the spectra. The spectra can be data as obtained in XRF, UV/VIS, Fluorescence, AES, NIR, IR, Raman, NMR, MS, etc. More generally, any data that is recorded over a discretized variable, e.g. absorbance = $f(\text{wavelength})$, stored as a vector of absorbance values for discrete wavelengths is suitable.

License GPL (≥ 3)

LazyLoad yes

LazyData yes

Depends R ($\geq 3.6.0$), lattice, grid, ggplot2 ($\geq 2.2.0$), xml2

Suggests R.matlab, deldir, rgl, plotrix, sp, baseline, compiler, inline, Repp, MASS, fastcluster, pls, mvtnorm, digest, reshape, devtools, R.rsp, tibble, knitr, rmarkdown

Imports testthat, methods, utils, latticeExtra, lazyeval, dplyr, rlang

URL <https://r-hyperspec.github.io/hyperSpec/> (documentation),
<https://github.com/r-hyperspec/hyperSpec> (code)

BugReports <https://github.com/r-hyperspec/hyperSpec/issues>

VignetteBuilder knitr

Collate 'validate.R' 'hyperspec-class.R' 'unittest.R' 'paste.row.R'
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 'write.txt.wide.R' 'y-pastenames.R' 'zzz.R'

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hyperSpec-package	<i>Package hyperSpec</i>
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Description

Interface for hyperspectral data sets This package gives an interface to handle hyperspectral data sets in R. Hyperspectral data are spatially or time-resolved spectra, or spectra with any other kind of information associated with the spectra. E.g. spectral maps or images, time series, calibration series, etc.

Details

The spectra can be data as obtained in XRF, UV/VIS, Fluorescence, AES, NIR, IR, Raman, NMR, MS, etc.

More generally, any data that is recorded over a discretized variable, e.g. absorbance = f (wavelength), stored as a vector of absorbance values for discrete wavelengths is suitable.

Author(s)

C. Beleites

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See Also

`citation("hyperSpec")` produces the correct citation.

`package?hyperSpec` for information about the package

`class?hyperSpec` for details on the S4 class provided by this package.

<code>.cluster.wavelengths</code>	<i>Find clusters of approximately equal wavelengths</i>
-----------------------------------	---

Description

Find clusters of approximately equal wavelengths

Usage

```
.cluster.wavelengths(dots, wl.tolerance)
```

Arguments

<code>dots</code>	list of hyperSpec objects to collapse
<code>wl.tolerance</code>	wavelength difference tolerance

Value

data.frame with information about suitable wavelength bins

.collapse.equal	<i>Try finding groups of hyperSpec objects with (approximately) equal wavelength axes</i>
-----------------	---

Description

... and directly rbind.fill them.

Usage

```
.collapse.equal(dots, wl.tolerance)
```

Arguments

dots	list with hyperSpec object to collapse
wl.tolerance	wavelength difference tolerance

Value

possible shorter list of dots

.DollarNames.hyperSpec	<i>command line completion for \$</i>
------------------------	---------------------------------------

Description

command line completion for \$

Usage

```
## S3 method for class 'hyperSpec'
.DollarNames(x, pattern = "")
```

Arguments

x	the hyperSpecobject
pattern	pattern to look for

Value

the name of the extra data slot

Author(s)

C. Beleites

See Also[.DollarNames](#)

<code>.fix_spc_colnames</code>	<i>Ensure that the spectra matrix has the wavelengths in column names</i>
--------------------------------	---

Description

Ensure that the spectra matrix has the wavelengths in column names

Usage`.fix_spc_colnames(spc)`**Arguments**

<code>spc</code>	hyperSpec object
------------------	------------------

ValuehyperSpec object with wavelengths in column names of `$spc`

<code>.read.spe.xml</code>	<i>Read XML footer from SPE file format version 3.0</i>
----------------------------	---

Description

The new SPE file format, introduced in 2012, was designed to be backwards compatible with the previous format 2.5. The most prominent change is the new plain text XML footer holding vast experimental metadata that gets attached at the end of the file. Thus, the file contains 3 blocks: a 4100-bytes long binary header, a chunk with spectral data, and the XML footer. This function retrieves the XML footer converted to R list, and throws error if it is not available. The file format specification is available at Princeton Instruments FTP server under name 'SPE 3.0 File Format Specification'.

Usage`.read.spe.xml(filename)`**Arguments**

<code>filename</code>	- SPE filename
-----------------------	----------------

Details

This function relies on R package `xml2` to work correctly

Value

xml data from the file converted to R list

```
.read.spe.xml_string    .read.spe.xml_string
```

Description

Read XML footer from SPE file format version 3.0 and return it as a long string for subsequent parsing. Basically the purpose of this function is to check that the file format version is 3.0 or above, and to find and read the correct part of this file.

Usage

```
.read.spe.xml_string(filename)
```

Arguments

`filename` - SPE filename

Value

string containing XML footer

```
aggregate                    aggregate hyperSpec objects
```

Description

Compute summary statistics for subsets of a `hyperSpec` object.

Usage

```
## S4 method for signature 'hyperSpec'
aggregate(
  x,
  by = stop("by is needed"),
  FUN = stop("FUN is needed."),
  ...,
  out.rows = NULL,
  append.rows = NULL,
  by.isindex = FALSE
)
```


Arguments

<code>x</code>	a hyperSpec object
<code>by</code>	grouping for the rows of <code>x@data</code> . Either a list containing an index vector for each of the subgroups or a vector that can be split in such a list.
<code>FUN</code>	function to compute the summary statistics
<code>...</code>	further arguments passed to <code>FUN</code>
<code>out.rows</code>	number of rows in the resulting hyperSpec object, for memory preallocation.
<code>append.rows</code>	If more rows are needed, how many should be appended? Defaults to 100 or an estimate based on the percentage of groups that are still to be done, whatever is larger.
<code>by.isindex</code>	If a list is given in <code>by</code> : does the list already contain the row indices of the groups? If <code>FALSE</code> , the list in <code>by</code> is computed first (as in aggregate).

Details

`aggregate` applies `FUN` to each of the subgroups given by `by`. It combines the functionality of [aggregate](#), [tapply](#), and [ave](#) for hyperSpec objects.

`aggregate` avoids splitting `x@data`.

`FUN` does not need to return exactly one value. The number of returned values needs to be the same for all wavelengths (otherwise the result could not be a matrix), see the examples.

If the initially preallocated `data.frame` turns out to be too small, more rows are appended and a warning is issued.

Value

A hyperSpec object with an additional column `@data$.aggregate` tracing which group the rows belong to.

Author(s)

C. Beleites

See Also

[tapply](#), [aggregate](#), [ave](#)

Examples

```
cluster.means <- aggregate (chondro, chondro$clusters, mean_pm_sd)
plot(cluster.means, stacked = ".aggregate", fill = ".aggregate",
      col = matlab.dark.palette (3))

## make some "spectra"
spc <- new ("hyperSpec", spc = sweep (rnorm (10*20), ncol = 20), 1, (1:10)*5, "+")

## 3 groups
```

```

color <- c("red", "blue", "black")
by <- as.factor (c (1, 1, 1, 1, 1, 1, 5, 1, 2, 2))
by
plot (spc, "spc", col = color[by])

## Example 1: plot the mean of the groups
plot (aggregate (spc, by, mean), "spc", col = color, add = TRUE,
      lines.args = list(lwd = 3, lty = 2))

## Example 2: FUN may return more than one value (here: 3)
plot (aggregate (spc, by, mean_pm_sd), "spc",
      col = rep(color, each = 3), lines.args = list(lwd = 3, lty = 2))

## Example 3: aggregate even takes FUN that return different numbers of
##           values for different groups
plot (spc, "spc", col = color[by])

weird.function <- function (x){
  if (length (x) == 1)
    x + 1 : 10
  else if (length (x) == 2)
    NULL
  else
    x [1]
}

agg <- aggregate (spc, by, weird.function)
agg$.aggregate
plot (agg, "spc", add = TRUE, col = color[agg$.aggregate],
      lines.args = list (lwd = 3, lty = 2))

```

apply

apply Computes summary statistics for the spectra of a hyperSpec object.

Description

apply gives the functionality of [apply](#) for hyperSpec objects.

Usage

```

## S4 method for signature 'hyperSpec'
apply(
  X,
  MARGIN,
  FUN,
  ...,
  label.wl = NULL,

```

```

    label.spc = NULL,
    new.wavelength = NULL,
    simplify
  )

```

Arguments

<code>X, spc</code>	a hyperSpec object
<code>MARGIN</code>	The subscript which the function will be applied over. 1 indicates rows (FUN is applied to each spectrum), 2 indicates columns (FUN is applied to each wavelength), 1 : 2 indicates that FUN should be applied to each single element of the spectra matrix. Note that many basic mathematical functions are already defined for hyperSpec objects (see Math). If MARGIN is missing, the whole spectra matrix is handed to FUN, see also the examples.
<code>FUN</code>	function to compute the summary statistics
<code>...</code>	further arguments passed to FUN
<code>label.wl, label.spc</code>	new labels for wavelength and spectral intensity axes
<code>new.wavelength</code>	for MARGIN = 2: numeric vector or name of the argument in ... that is to be used (character) as wavelength axis of the resulting object.
<code>simplify</code>	ignored: apply for hyperSpec results are always simplified

Details

The generic functions of group [Math](#) are not defined for hyperSpec objects. Instead, `apply` can be used. For functions like `log` that work on scalars, `MARGIN = 1 : 2` gives the appropriate behaviour.

`spcapply` does the same as `apply` with `MARGIN = 1`, but additionally allows to set a new wavelength axis and adjust the labels.

`wlapply` does the same as `apply` with `MARGIN = 2`, but additionally allows to set a new wavelength axis and adjust the labels.

Value

A hyperSpec object

Author(s)

C. Beleites

See Also

[apply](#), for applying FUN to subgroups of the hyperSpec object: [aggregate](#).

Examples

```

plotspc (apply (chondro, 2, range))

avgflu <- apply (flu, 1, mean,
                label.spc = expression (bar (I)),
                new.wavelength = mean (wl (flu)))
avgflu

flu[, ,405:407]
apply (flu, 1:2, "*", -1)[, ,405:407]

## without MARGIN the whole matrix is handed to FUN
apply (flu [, ,405:407], , print) [[]]

## whereas MARGIN = 1 : 2 leads to FUN being called for each element separately
apply (flu [, ,405:407], 1 : 2, print) [[]]

```

Arith

Arithmetical Operators for hyperSpec objects

Description

Arithmetical Operators: +, -, *, /, ^, %%, %!%, %*% for hyperSpec objects

Usage

```

## S4 method for signature 'hyperSpec,hyperSpec'
Arith(e1, e2)

## S4 method for signature 'hyperSpec,numeric'
Arith(e1, e2)

## S4 method for signature 'hyperSpec,matrix'
Arith(e1, e2)

## S4 method for signature 'hyperSpec,missing'
Arith(e1, e2)

## S4 method for signature 'numeric,hyperSpec'
Arith(e1, e2)

## S4 method for signature 'matrix,hyperSpec'
Arith(e1, e2)

## S4 method for signature 'hyperSpec,hyperSpec'
x %*% y

```

```
## S4 method for signature 'hyperSpec,matrix'
x %**% y

## S4 method for signature 'matrix,hyperSpec'
x %**% y
```

Arguments

e1, e2 or
 x, y either two hyperSpec objects or
 one hyperSpec object and matrix of same size as hyperSpec[[]] or
 a vector which length equalling either the number of rows or the number of
 wavelengths of the hyperSpec object, or
 a scalar (numeric of length 1).

Details

The arithmetical operators +, -, *, /, ^, %, %/, and %**% for hyperSpec objects.

You can use these operators in different ways:

```
e1 + e2
`+` (e1, e2)
```

```
x %**% y
`%**%`(x, y)
```

```
-x
```

The arithmetical operators +, -, *, /, ^, %, %/, and %**% work on the spectra matrix of the hyperSpec object. They have their usual meaning (see [Arithmetic](#)). The operators work also with one hyperSpec object and a numeric object or a matrices of the same size as the spectra matrix of the hyperSpec object.

With numeric vectors [sweep](#) is most probably more appropriate.

If you want to calculate on the extra data as well, use the data.frame hyperSpec@data directly or [as.data.frame](#) (x).

Value

hyperSpec object with the new spectra matrix.

Author(s)

C. Beleites

See Also

[sweep-methods](#) for calculations involving a vector and the spectral matrix.

[S4groupGeneric](#) for group generic methods.

[Arithmetic](#) for the base arithmetic functions.

[Comparison](#) for comparison operators, [Math](#) for mathematical group generic functions (Math and Math2 groups) working on hyperSpec objects.

[matmult](#) for matrix multiplications with %*%.

Examples

```
flu + flu
1 / flu
all((flu + flu - 2 * flu)[][] == 0)
-flu
flu / flu$c
```

as.character,hyperSpec-method

Convert a hyperSpec object to character strings for Display print, show, and summary show the result of as.character.

Description

print, show, and summary differ only in the defaults. show displays the range of values instead,

Usage

```
## S4 method for signature 'hyperSpec'
as.character(
  x,
  digits = getOption("digits"),
  range = TRUE,
  max.print = 5,
  shorten.to = c(2, 1)
)

## S4 method for signature 'hyperSpec'
show(object)

## S4 method for signature 'hyperSpec'
print(x, range = FALSE, ...)

## S4 method for signature 'hyperSpec'
summary(object, ...)
```

Arguments

x	a hyperSpec object
digits	number of digits handed over to format
range	should the values be indicated as range rather than first and last elements?
max.print	maximum number of elements to be printed (of a variable)
shorten.to	if a vector is longer than max.print, only the first shorten.to[1] and the last shorten.to[2] elements are printed
object	a hyperSpec object
...	print and summary hand further arguments to as.character

Value

as.character returns a character vector fit to be printed by cat with sep = "\n".
 print invisibly returns x after printing, show returns an invisible NULL.

See Also

[as.character](#)
[show](#)
[print](#)
[summary](#)

Examples

```
chondro

show (chondro)

summary (chondro)

print (chondro, range = TRUE)
```

as.data.frame	<i>Conversion of a hyperSpec object into a data.frame or matrix as.data.frame returns x@data (as data.frame) as.matrix returns the spectra matrix x@data\$spc as matrix</i>
---------------	---

Description

The data.frame returned by as.long.df is guaranteed to have columns spc and .wavelength. If nwl (x) == 0 these columns will be NA.

Usage

```
## S3 method for class 'hyperSpec'
as.data.frame(x, row.names = TRUE, optional = NULL, ...)

## S3 method for class 'hyperSpec'
as.matrix(x, ...)

as.wide.df(x, wl.prefix = "")

as.long.df(x, rownames = FALSE, wl.factor = FALSE, na.rm = TRUE)

as.t.df(x)
```

Arguments

x	a hyperSpec object
row.names	if TRUE, a column .row is created containing row names or row indices if no rownames are set. If character vector, the rownames are set accordingly.
optional	ignored
...	ignored
wl.prefix	prefix to prepend wavelength column names
rownames	should the rownames be in column .rownames of the long-format data.frame?
wl.factor	should the wavelengths be returned as a factor (instead of numeric)?
na.rm	if TRUE, rows where spc is not NA are deleted.

Value

x@data and x@data\$spc (== x\$spc == x [[]]), respectively.

as.wide.df returns a data.frame that consists of the extra data and the spectra matrix converted to a data.frame. The spectra matrix is expanded *in place*.

as.long.df returns the stacked or molten version of x@data. The wavelengths are in column .wavelength.

as.t.df returns a data.frame similar to as.long.df, but each spectrum in its own column. This is useful for exporting summary spectra, see the example.

Author(s)

C. Beleites

See Also

[as.data.frame](#)
 and `base::as.matrix()`
`[[[]]]` (`[[[]]]`) for a shortcut to `as.matrix`
[stack](#) and [melt](#) or `reshape2::melt()` for other functions producing long-format data.frames.

Examples

```

as.data.frame (chondro [1:3,, 600 ~ 620])
as.matrix (chondro [1:3,, 600 ~ 620])
lm (c ~ spc, data = flu [,450])

as.wide.df (chondro [1:5,, 600 ~ 610])
summary (as.wide.df (chondro [1:5,, 600 ~ 610]))

as.long.df (flu [, 405 ~ 410])
summary (as.long.df (flu [, 405 ~ 410]))
summary (as.long.df (flu [, 405 ~ 410], rownames = TRUE))
summary (as.long.df (flu [, 405 ~ 410], wl.factor = TRUE))

df <- as.t.df (apply (chondro, 2, mean_pm_sd))
head (df)

if (require (ggplot2)){
  ggplot (df, aes (x = .wavelength)) +
    geom_ribbon (aes (ymin = mean.minus.sd, ymax = mean.plus.sd),
      fill = "#00000040") +
    geom_line (aes (y = mean))
}

```

as.hyperSpec

*as.hyperSpec: convenience conversion functions***Description**

These functions are shortcuts to convert other objects into hypeSpec objects.

Usage

```

as.hyperSpec(X, ...)

## S4 method for signature 'matrix'
as.hyperSpec(X, wl = guess.wavelength(colnames(X)), ...)

## S4 method for signature 'data.frame'
as.hyperSpec(
  X,
  spc = NULL,
  wl = guess.wavelength(spc),
  labels = attr(X, "labels"),
  ...
)

```

Arguments

<code>X</code>	the object to convert. A matrix is assumed to contain the spectra matrix, a data.frame is assumed to contain extra data.
<code>...</code>	additional parameters that should be handed over to new ("hyperSpec") (initialize)
<code>wl</code>	wavelength vector. Defaults to guessing from the column names in <code>X</code>
<code>spc</code>	spectra matrix
<code>labels</code>	list with labels

Value

hyperSpec object

Note

Note that the behaviour of `as.hyperSpec(X)` was changed: it now assumes `X` to be extra data, and returns a hyperSpec object with 0 wavelengths. To get the old behaviour

See Also

[initialize](#)

Examples

```
tmp <- data.frame(flu[, , 400 ~ 410])
(wl <- colnames(tmp))
guess.wavelength(wl)
```

barbiturates

Barbiturates Spectra from .spc example files A time series of mass spectra in a list of hyperSpec objects.

Description

Barbiturates Spectra from .spc example files A time series of mass spectra in a list of hyperSpec objects.

Format

The data sets consists of 286 spectra. They are the result of importing the BARBITUATES.SPC example data from Thermo Galactic's spc file format specification.

Author(s)

C. Beleites and Thermo Galactic

References

The raw data is available at <http://hyperspec.r-forge.r-project.org/blob/fileio.zip>

Examples

```
barbiturates [1:3]
length (barbiturates)

barb <- collapse (barbiturates, collapse.equal = FALSE)
barb <- orderwl (barb)

plot (barb [1:3], lines.args = list (type = "h"),
      col = matlab.dark.palette (3), stacked = TRUE,
      stacked.args = list (add.factor = .2))

if (require (latticeExtra)){
  levelplot (spc ~ .wavelength * z, log (barb), panel = panel.levelplot.points,
            cex = 0.3, col = "#00000000", col.regions = matlab.palette (20))
}

plotc (apply (barb [, 42.9~43.2], 1, sum, na.rm = TRUE), spc ~ z,
       panel = panel.lines, ylab = expression (I[m/z == 43] / "a.u."))
```

bind

Binding hyperSpec Objects

Description

The former difficulties with binding S4 objects are resolved since R version 3.2.0 and cbind and rbind now work as intended and expected for hyperSpec objects.

cbind2 binds the spectral matrices of two hyperSpec objects by column. All columns besides spc with the same name in x@data and y@data must have the same elements. Rows are ordered before checking.

Usage

```
bind(
  direction = stop("direction ('c' or 'r') required"),
  ...,
  wl.tolerance = hy.getOption("wl.tolerance")
)

## S3 method for class 'hyperSpec'
cbind(...)

## S3 method for class 'hyperSpec'
rbind(...)
```

```
## S4 method for signature 'hyperSpec,hyperSpec'
cbind2(x, y)

## S4 method for signature 'hyperSpec,missing'
cbind2(x, y)

## S4 method for signature 'hyperSpec,hyperSpec'
rbind2(x, y, wl.tolerance = hy.getOption("wl.tolerance"))

## S4 method for signature 'hyperSpec,missing'
rbind2(x, y, wl.tolerance)
```

Arguments

direction	"r" or "c" to bind rows or columns
...	The hyperSpec objects to be combined. Alternatively, <i>one</i> list of hyperSpec objects can be given to bind.
wl.tolerance	rbind and rbind2 check for equal wavelengths with this tolerance.
x, y	hyperSpec objects

Details

Therefore, calling `rbind.hyperSpec` and `cbind.hyperSpec` is now deprecated: `cbind` and `rbind` should now be called directly.

However, in consequence it is no longer possible to call `cbind` or `rbind` with a list of `hyperSpec` objects. In that case, use `bind` or [do.call](#) (see example).

`bind` does the common work for both column- and row-wise binding.

Value

a `hyperSpec` object, possibly with different row order (for `bind("c", ...{ })` and `cbind2`).

Note

You might have to make sure that the objects either all have or all do not have rownames and/or colnames.

Author(s)

C. Beleites

See Also

[rbind2](#), [cbind2](#) [rbind](#), [cbind](#)

[merge](#) and [collapse](#) for combining objects that do not share spectra or wavelengths, respectively.

Examples

```

chondro
bind ("r", chondro, chondro)
rbind (chondro, chondro)
cbind (chondro, chondro)
bind ("r", list (chondro, chondro, chondro))

x <- chondro[, 600 : 605]
x$a <- 1
x@data <- x@data[, sample (ncol (x), ncol (x))] # reorder columns

y <- chondro [nrow (chondro) : 1,, 1730 : 1750] # reorder rows
y$b <- 2

cbind2 (x, y) # works

y$y[3] <- 5
try (cbind2 (x, y)) # error

# list of hyperSpec objects

lhy <- list (flu, flu)
do.call ("rbind", lhy)
bind ("r", lhy)

```

chk.hy

*Validation of hyperSpec objects***Description**

Check whether an object is a hyperSpec object and validate the object

Usage

```
chk.hy(object)
```

Arguments

object the object to check

Value

TRUE if the check passes, otherwise stop with an error.

Author(s)

C. Beleites

See Also`validObject`**Examples**

```
chk.hy (chondro)
validObject (chondro)
```

chondro

Raman spectra of 2 Chondrocytes in Cartilage A Raman-map (laterally resolved Raman spectra) of chondrocytes in cartilage.

Description

See the vignette `vignette("chondro", package = "hyperSpec")`.

Usage

```
chondro
```

Format

The data set has 875 Raman spectra measured on a 25×35 grid with 1 micron step size. Spatial information is in `chondro$x` and `chondro$y`. Each spectrum has 300 data points in the range of ca. 600 - 1800 cm^{-1} .

Author(s)

A. Bonifacio and C. Beleites

References

The raw data is available at <http://hyperspec.r-forge.r-project.org/blob/chondro.zip>

Examples

```
chondro

## do baseline correction
baselines <- spc.fit.poly.below (chondro)
chondro <- chondro - baselines

## area normalization
chondro <- chondro / colMeans (chondro)

## subtract common composition
chondro <- chondro - quantile (chondro, 0.05)
```

```

cols <- c("dark blue", "orange", "#C02020")
plotmap (chondro, clusters ~ x * y, col.regions = cols)

cluster.means <- aggregate (chondro, chondro$clusters, mean_pm_sd)
plot (cluster.means, stacked = ".aggregate", fill = ".aggregate", col = cols)

## plot nucleic acid bands
plotmap (chondro[, , c( 728, 782, 1098, 1240, 1482, 1577)],
        col.regions = colorRampPalette (c ("white", "gold", "dark green"), space = "Lab") (20))

```

collapse

*Collapse hyperSpec objects***Description**

collapse/bind several hyperSpec objects into one object

Usage

```

collapse(
  ...,
  wl.tolerance = hy.getOption("wl.tolerance"),
  collapse.equal = TRUE
)

```

Arguments

... hyperSpec objects to be collapsed into one object. Instead of giving several arguments, a list with all objects to be collapsed may be given.

wl.tolerance tolerance to decide which wavelengths are considered equal.

collapse.equal logical indicating whether to try first finding groups of spectra with (approximately) equal wavelength axes. If the data is known to contain few or no such groups, collapse() will be faster with this first pass being turned off.

Details

The spectra from all objects will be put into one object. The resulting object has all wavelengths that occur in any of the input objects, wl.tolerance is used to determine which difference in the wavelengths is tolerated as equal: clusters of approximately equal wavelengths will span at most $2 * \text{wl.tolerance}$. Differences up to $\pm \text{wl.tolerance}$ are considered equal.

The returned object has wavelengths that are the weighted average (by number of spectra) of the wavelengths within any such cluster of approximately equal wavelengths.

Labels will be taken from the first object where they are encountered. However, the order of processing objects is not necessarily the same as the order of objects in the input: collapse first processes groups of input objects that share all wavelengths (within wl.tolerance).

Data points corresponding to wavelengths not in the original spectrum will be set to NA. Extra data is combined in the same manner.

If the objects are named, the names will be preserved in extra data column \$.name. If the wavelengths are names, names are preserved and taken from the first object where they were encountered, the same applies to possible column names of the spectra matrix.

Value

a hyperSpec object

Author(s)

C. Beleites

See Also

[merge\(\)](#), [rbind\(\)](#), and [plyr::rbind.fill\(\)](#)

Examples

```
barbiturates [1:3]
collapse (barbiturates [1:3])

a <- barbiturates [[1]]
b <- barbiturates [[2]]
c <- barbiturates [[3]]

a
b
c
collapse (a, b, c)

collapse (barbiturates [1:3], collapse.equal = FALSE)
```

colSums

colSums, colMeans, rowSums and rowMeans functions for hyperSpec objects

Description

hyperSpec objects can use the base functions [colMeans](#), [colSums](#), [rowMeans](#) and [rowSums](#).

Usage

```
## S4 method for signature 'hyperSpec'
colMeans(x, na.rm = TRUE, ..., label.spc)

## S4 method for signature 'hyperSpec'
colSums(x, na.rm = TRUE, ..., label.spc)

## S4 method for signature 'hyperSpec'
rowMeans(x, na.rm = TRUE, ..., label.wavelength)

## S4 method for signature 'hyperSpec'
rowSums(x, na.rm = TRUE, ..., label.wavelength)
```

Arguments

<code>x</code>	hyperSpec object
<code>na.rm, ...</code>	further parameters to the base functions na.rm defaults to TRUE for hyperSpec objects.
<code>label.spc</code>	labels for the intensity axis for loadings-like (col) statistics
<code>label.wavelength</code>	labels for the wavelength axis for scores-like (row) statistics

See Also

[colSums](#)

Examples

```
colMeans (flu)
colSums (flu)
colSums (flu)
rowSums (flu)
```

Comparison

Comparison of hyperSpec objects

Description

The comparison operators `>`, `<`, `>=`, `<=`, `==`, and `!=` for hyperSpec objects.

Usage

```
## S4 method for signature 'hyperSpec,hyperSpec'
Compare(e1, e2)

## S4 method for signature 'hyperSpec,numeric'
```

```

Compare(e1, e2)

## S4 method for signature 'hyperSpec,matrix'
Compare(e1, e2)

## S4 method for signature 'numeric,hyperSpec'
Compare(e1, e2)

## S4 method for signature 'matrix,hyperSpec'
Compare(e1, e2)

## S4 method for signature 'hyperSpec,hyperSpec'
all.equal(
  target,
  current,
  ...,
  check.attributes = FALSE,
  check.names = FALSE,
  check.column.order = FALSE,
  check.label = FALSE,
  tolerance = hy.getOption("tolerance"),
  wl.tolerance = hy.getOption("wl.tolerance")
)

```

Arguments

e1, e2	Either two hyperSpec objects or one hyperSpec object and matrix of same size as hyperSpec[[]] or a scalar (numeric of length 1). As hyperSpec objects must have numeric spectra matrices, the resulting matrix of the comparison is returned directly.
target, current	two hyperSpec objects that are tested for equality
...	handed to all.equal when testing the slots of the hyperSpec objects
check.attributes, check.names	see all.equal
check.column.order	If two objects have the same data, but the order of the columns (determined by the names) differs, should they be regarded as different?
check.label	Should the slot label be checked? If the labels differ only in the order of their entries, they are considered equal.
tolerance, wl.tolerance	tolerances for checking wavelengths and data, respectively

Details

`all.equal` checks the equality of two hyperSpec objects.

The comparison operators `>`, `<`, `>=`, `<=`, `==`, and `!=` work on the spectra matrix of the hyperSpec object. They have their usual meaning (see [Comparison](#)). The operators work also with one

hyperSpec object and a numeric (scalar) object or a matrices of the same size as the spectra matrix of the hyperSpec object.

With numeric vectors [sweep](#) might be more appropriate.

If you want to calculate on the data.frame hyperSpec@data, you have to do this directly on hyperSpec@data.

Value

a logical matrix for the comparison operators.

`all.equal` returns either TRUE, or a character vector describing the differences. In conditions, the result must therefore be tested with [isTRUE](#).

Author(s)

C. Beleites

See Also

[sweep-methods](#) for calculations involving a vector and the spectral matrix.

[S4groupGeneric](#) for group generic methods.

[Comparison](#) for the base comparison functions.

[Arith](#) for arithmetic operators, [Math](#) for mathematical group generic functions (groups Math and Math2) working on hyperSpec objects.

[all.equal](#) and [isTRUE](#)

Examples

```
flu[, , 445 ~ 450] > 300
```

```
all (flu == flu[[ ]])
```

count_lines	<i>count lines (of an ASCII file)</i>
-------------	---------------------------------------

Description

count lines (of an ASCII file)

Usage

```
count_lines(file, chunksize = 10000)
```

Arguments

file	the file name or connection
chunksize	file is read in chunks of chunksize lines.

Value

number of lines in file

Author(s)

C. Beleites

cov,hyperSpec,missing-method

Covariance matrices for hyperSpec objects

Description

Covariance matrices for hyperSpec objects

Usage

```
## S4 method for signature 'hyperSpec,missing'
cov(
  x,
  y = NULL,
  use = "everything",
  method = c("pearson", "kendall", "spearman")
)
```

```
pooled.cov(x, groups, ..., regularize = 1e-05 * max(abs(COV)))
```

Arguments

x	hyperSpec object
y	not supported
use, method	handed to cov
groups	factor indicating the groups
...	ignored
regularize	regularization of the covariance matrix. Set 0 to switch off pooled.cov calculates pooled covariance like e.g. in LDA.

Value

covariance matrix of size `nwl(x) x nwl(x)`

Author(s)

C. Beleites

See Also[cov](#)**Examples**

```
image (cov (chondro))
pcov <- pooled.cov (chondro, chondro$clusters)
plot (pcov$means)
image (pcov$COV)
```

decomposition	<i>Convert Principal Component Decomposition or the like into a hyperSpec Object</i>
---------------	--

Description

Decomposition of the spectra matrix is a common procedure in chemometric data analysis. scores and loadings convert the result matrices into new hyperSpec objects.

Usage

```
decomposition(
  object,
  x,
  wavelength = seq_len(ncol(x)),
  label.wavelength,
  label.spc,
  scores = TRUE,
  retain.columns = FALSE,
  ...
)
```

Arguments

object	A hyperSpec object.
x	matrix with the new content for object@data\$spc. Its size must correspond to rows (for scores) and to either columns or rows (for loadings) of object.
wavelength	for a scores-like x: the new object@wavelength.
label.wavelength	The new label for the wavelength axis (if x is scores-like). If not given, the label of object is kept.
label.spc	The new label for the spectra matrix. If not given, the label of object is kept.
scores	is x a scores-like matrix?

`retain.columns` for loading-like decomposition (i.e. `x` holds loadings, pure component spectra or the like), the data columns need special attention. Columns with different values across the rows will be set to NA if `retain.columns` is TRUE, otherwise they will be deleted.

... ignored.

Details

Multivariate data are frequently decomposed by methods like principal component analysis, partial least squares, linear discriminant analysis, and the like. These methods yield latent spectra (or latent variables, loadings, components, ...) that are linear combination coefficients along the wavelength axis and scores for each spectrum and loading.

The loadings matrix gives a coordinate transformation, and the scores are values in that new coordinate system.

The obtained latent variables are spectra-like objects: a latent variable has a coefficient for each wavelength. If such a matrix (with the same number of columns as object has wavelengths) is given to `decomposition` (also setting `scores = FALSE`), the spectra matrix is replaced by `x`. Moreover, all columns of `object@data` that did not contain the same value for all spectra are set to NA. Thus, for the resulting `hyperSpec` object, `plotspc` and related functions are meaningful. `plotmap` cannot be applied as the loadings are not laterally resolved.

The scores matrix needs to have the same number of rows as object has spectra. If such a matrix is given, `decomposition` will replace the spectra matrix is replaced by `x` and `object@wavelength` by `wavelength`. The information related to each of the spectra is retained. For such a `hyperSpec` object, `plotmap` and `plotc` and the like can be applied. It is also possible to use the spectra plotting, but the interpretation is not that of the spectrum any longer.

Value

A `hyperSpec` object, updated according to `x`

Author(s)

C. Beleites

See Also

See `%*%` for matrix multiplication of `hyperSpec` objects.

See e.g. `prcomp` and `princomp` for principal component analysis, and package `pls` for Partial Least Squares Regression.

Examples

```
pca <- prcomp (flu)

pca.loadings <- decomposition (flu, t (pca$rotation), scores = FALSE)
pca.center <- decomposition (flu, pca$center, scores = FALSE)
pca.scores <- decomposition (flu, pca$x)
```

```
plot (pca.center)
plot (pca.loadings, col = c ("red", "gray50"))
plotc (pca.scores, groups = .wavelength)
```

dimnames,hyperSpec-method

dimnames for hyperSpec objects

Description

hyperSpec objects can have row- and column names like data.frames. The "names" of the wavelengths are treated separately: see [wl](#)

Usage

```
## S4 method for signature 'hyperSpec'
dimnames(x)

## S4 method for signature 'hyperSpec'
rownames(x, do.NULL = TRUE, prefix = "row")

## S4 replacement method for signature 'hyperSpec'
rownames(x) <- value

## S4 method for signature 'hyperSpec'
colnames(x, do.NULL = TRUE, prefix = "col")

## S4 replacement method for signature 'hyperSpec'
colnames(x) <- value
```

Arguments

x	the hyperSpec object
do.NULL	handed to rownames or colnames : logical. Should this create names if they are NULL?
prefix	handed to rownames or colnames
value	the new names

Author(s)

C. Beleites

See Also

[wl](#) for the wavelength dimension

[dimnames](#)

[rownames](#)

[colnames](#)

Examples

```
dimnames (flu)
rownames (flu)
colnames (chondro)
```

droplevels,hyperSpec-method
droplevels for hyperSpec object

Description

calls `base::droplevels()` on the data.frame in `spc@data`.

Usage

```
## S4 method for signature 'hyperSpec'
droplevels(x, ...)
```

Arguments

x	hyperSpec object
...	handed to <code>base::droplevels.data.frame()</code>

Value

hyperSpec object with unused levels of all factors in @data dropped.

See Also

`base::droplevels()`

Examples

```
chondro[1:3]$clusters
droplevels (chondro [1:3])$clusters
```

empty	<i>Empty hyperSpec object</i>
-------	-------------------------------

Description

Empty produces an hyperSpec object with the same columns and wavelengths as x. The new object will either contain no rows at all (default), or the given number of rows with all data initialized to spc and extra, respectively.

Usage

```
empty(x, nrow = 0, spc = NA, extra = NA)
```

Arguments

x	hyperSpec object
nrow	number of rows the new object should have
spc	value to initialize the new spectra matrix with
extra	value to initialize the new extra data with

Author(s)

C. Beleites

Examples

```
empty (chondro, nrow = 2, spc = 0)
```

flu	<i>Quinine Fluorescence Spectra Fluorescence spectra of different dilutions of quinine forming a calibration set.</i>
-----	---

Description

See the vignette: vignette ("flu", package = "hyperSpec")

Format

The data set has 6 fluorescence emission spectra measured on quinine concentrations between 0.05 mg/l and 0.30 mg/l. Each spectrum consists of 181 data points in the range of 405 nm to 495 nm.

Author(s)

M. Kammer and C. Beleites

Examples

```
flu
plot (flu)
plotc (flu)
```

Future-functions	<i>Future functions</i>
------------------	-------------------------

Description

These functions will be introduced in **hyperSpec** v1.0 and will replace some current functions. Now they appear here just for compatibility with other packages, which should be released on CRAN. They are not intended to be used by **hyperSpec** v0.100 users directly.

Usage

```
.spc_io_postprocess_optional(...)
wl_convert_units(x, from, to, ref_wl = NULL)
hy_set_options(...)
hy_get_option(...)
hy_set_options(...)
read_txt_long(...)
read_txt_wide(...)
.wl_fix_unit_name(...)
assert_hyperSpec(...)
```

Arguments

```
...           Arguments to functions.
x, from, to, ref_wl
              Arguments to functions.
```

guess.wavelength	<i>guess wavelengths from character vector</i>
------------------	--

Description

character vectors used for names (e.g. colnames for matrices or data.frames) are often treated by [make.names](#) or similar functions that produce suitable names (e.g. by prepending "X" to numbers.). Such names cannot be directly converted to numeric.

Usage

```
guess.wavelength(X)
```

Arguments

X character with numbers hidden inside

Details

guess.wavelength tries to extract numbers from X which may be surrounded by such "protecting" characters.

Value

numeric

Examples

```
tmp <- data.frame(flu [[, 400 ~ 410]])
(wl <- colnames (tmp))
guess.wavelength (wl)
```

hy.getOptions	<i>Options for package hyperSpec Functions to access and set hyperSpec's options.</i>
---------------	---

Description

Currently, the following options are defined:

Name	Default Value (range)	Description
debuglevel	0 (1L 2L 3L)	amount of debugging information produced
gc	FALSE	triggers frequent calling of gc ()
file.remove.emptyspc	TRUE	remove empty spectra directly on file import

Used by
[spc.identify](#)
various file imp
[spc.fit.poly](#)
[read.ENVI](#), ne
various file imp

file.keep.name	TRUE	always create filename column	various file imp
tolerance	<code>sqrt (.Machine\$double.eps)</code>	tolerance for numerical comparisons	<code>normalize01</code> ,
wl.tolerance	<code>sqrt (.Machine\$double.eps)</code>	tolerance for comparisons of the wavelength axis	<code>all.equal</code> , <code>co</code>
plot.spc.nmax	25	number of spectra to be plotted by default	<code>plotspc</code>
ggplot.spc.nmax	10		<code>qplotspc</code>

Usage

```
hy.getOptions(...)  
  
hy.getOption(name)  
  
hy.setOptions(...)
```

Arguments

...	hy.setOptions: pairs of argument names and values. hy.getOptions: indices (or names) of the options.
name	the name of the option

Details

hy.setOptions will discard any values that were given without a name.

Value

hy.getOptions	returns a list of all options
hy.setOptions	invisibly returns a list with the options
hy.getOption	returns the value of the requested option

Author(s)

C. Beleites

Examples

```
hy.getOptions ()
```

hy.unittest	<i>hyperSpec unit tests</i>
-------------	-----------------------------

Description

If `testthat` is available, run the unit tests and display the results.

Usage

```
hy.unittest(standalone = TRUE, reporter = "progress")
```

Arguments

<code>standalone</code>	run the unit test on their own, e.g. from the console ('TRUE') or within testthat tests ('FALSE'), e.g. via 'devtools::test()'
<code>reporter</code>	the reporter to use, defaults to [testthat::ProgressReporter]

Value

Invisibly returns a data frame with the test results

Author(s)

Claudia Beleites

Examples

```
hy.unittest ()
```

hyperSpec-class	<i>Class "hyperSpec" This class handles hyperspectral data sets, i.e. spatially or time-resolved spectra, or spectra with any other kind of information associated with the spectra.</i>
-----------------	--

Description

The spectra can be data as obtained in XRF, UV/VIS, Fluorescence, AES, NIR, IR, Raman, NMR, MS, etc.

Details

More generally, any data that is recorded over a discretized variable, e.g. absorbance = f (wavelength), stored as a vector of absorbance values for discrete wavelengths is suitable.

Slots

`wavelength` wavelengths (wavenumbers, frequencies, etc.) for each of the columns of the spectra matrix

`data` the data (extra data and spectra matrix)

`label` expressions for column labels (incl. units). The label of the wavelength axis is in the special element `.wavelength`.

`log` deprecated.

Note

Please note that the logbook is now removed.

Author(s)

C. Beleites

See Also

See the vignette "hyperspec" for an introduction to hyperSpec from a spectroscopic point of view.

Examples

```
showClass("hyperSpec")
## Not run: vignette("hyperspec")
```

initialize

Creating a hyperSpec Object

Description

Like other S4 objects, a `hyperSpec` object can be created by `new`. The `hyperSpec` object is then initialized using the given parameters.

Usage

```
## S4 method for signature 'hyperSpec'
initialize(.Object, spc = NULL, data = NULL, wavelength = NULL, labels = NULL)
```

Arguments

`.Object` the new `hyperSpec` object.

`spc` the spectra matrix.
`spc` does not need to be a matrix, it is converted explicitly by `I(as.matrix(spc))`.

data	data.frame, possibly with the spectra in data\$spc, and further variates in more columns. A matrix can be entered as <i>one</i> column of a data frame by: <code>data.frame (spc = I (as.matrix (spc)))</code> . However, it will usually be more convenient if the spectra are given in spc
wavelength	The wavelengths corresponding to the columns of data. If no wavelengths are given, an appropriate vector is derived from the column names of data\$spc. If this is not possible, <code>1 : ncol (data\$spc)</code> is used instead.
labels	A list containing the labels for the columns of the data slot of the hyperSpec object and for the wavelength (in label\$.wavelength). The labels should be given in a form ready for the text-drawing functions (see plotmath). If label is not given, a list containing NULL for each of the columns of data and wavelength is used.

Details

If option gc is TRUE, the initialization will have frequent calls to `gc ()` which can help to avoid swapping or running out of memory.

Author(s)

C.Beleites

See Also

[new](#) for more information on creating and initializing S4 objects.

[plotmath](#) on expressions for math annotations as for slot label.

[hy.setOptions](#)

Examples

```
new ("hyperSpec")

spc <- matrix (rnorm (12), ncol = 4)
new ("hyperSpec", spc = spc)
new ("hyperSpec", data = data.frame (x = letters[1:3]),
    spc = spc)

colnames (spc) <- 600:603
new ("hyperSpec", spc = spc) # wavelength taken from colnames (spc)

# given wavelengths precede over colnames of spc
new ("hyperSpec", spc = spc, wavelength = 700:703)

# specifying labels
h <- new ("hyperSpec", spc = spc, data = data.frame (pos = 1 : 3),
    label = list (spc = "I / a.u.",
        .wavelength = expression (tilde (nu) / cm^-1),
        pos = expression ("/" (x, mu*m)))
)
```

```
plot (h)
plotc (h, spc ~ pos)
```

labels<-	<i>Get and Set Labels of a hyperSpec Object value may be a list or vector of labels giving the new label for each of the entries specified by which.</i>
----------	--

Description

The names of the labels are the same as the colnames of the data.frame. The label for the wavelength axis has the name .wavelength.

Usage

```
labels (object, which = NULL, ...) <- value

## S4 method for signature 'hyperSpec'
labels(object, which = bquote(), drop = TRUE, ..., use.colnames = TRUE)
```

Arguments

object	a hyperSpec object
which	numeric or character to specify the label(s)
...	ignored
value	the new label(s)
drop	if the result would be a list with only one element, should the element be returned instead?
use.colnames	should missing labels be replaced by column names of the extra data?

Details

The labels should be given in a form ready for the text-drawing functions (see [plotmath](#)), e.g. as expression or a character.

Value

labels<- returns a hyperSpec object.

labels returns a list of labels. If drop is TRUE and the list contains only one element, the element is returned instead.

Author(s)

C. Beleites

See Also[labels](#)**Examples**

```
labels (flu, "c") <- expression ("/" ("c", "mg / l"))
```

```
labels (chondro)
```

laser

Laser Emission A time series of an unstable laser emission.

Description

see the Vignette

Format

The data set consists of 84 laser emission spectra measured during 95 min. Each spectrum has 36 data points in the range of 404.5 nm to 405.8 nm.

Author(s)

C. Beleites

Examples

```
laser

cols <- c ("black", "blue", "darkgreen", "red")
wl <- c (405.0, 405.1, 405.3, 405.4)
plotspc (laser, axis.args=list (x = list (at = seq (404.5, 405.8, .1))))
for (i in seq_along (wl))
  abline (v = wl[i], col = cols[i], lwd = 2, lty = 2)

plotc (laser [, wl], spc ~ t, groups = .wavelength, type = "b",
       col = cols)

## Not run: vignette ("laser", package="hyperSpec")
```

legendright

Plot multivariate data into colour channels

Description

plot graph with legend right of it

Usage

```
legendright(p, l, legend.width = 8, legend.unit = "lines")

qmixtile(
  object,
  purecol = stop("pure component colors needed."),
  mapping = aes_string(x = "x", y = "y", fill = "spc"),
  ...,
  map.tileonly = FALSE
)

normalize.colrange(x, na.rm = TRUE, legend = FALSE, n = 100, ...)

normalize.range(x, na.rm = TRUE, legend = FALSE, n = 100, ...)

normalize.null(x, na.rm = TRUE, legend = FALSE, n = 100, ...)

normalize.minmax(x, min = 0, max = 1, legend = FALSE, n = 100, ...)

qmixlegend(
  x,
  purecol,
  dx = 0.33,
  ny = 100,
  labels = names(purecol),
  normalize = normalize.colrange,
  ...
)

colmix.rgb(
  x,
  purecol,
  against = 1,
  sub = TRUE,
  normalize = normalize.colrange,
  ...
)
```

Arguments

<code>p</code>	plot object
<code>l</code>	legend object
<code>legend.width, legend.unit</code>	size of legend part
<code>object</code>	matrix to be plotted with mixed colour channels
<code>purecol</code>	pure component colours, names determine legend labels
<code>mapping</code>	see geom_tile
<code>...</code>	qmixtile: handed to colmix.rgb qmixlegend and colmix.rgb hand further arguments to the normalize function
<code>map.tileonly</code>	if TRUE, mapping will be handed to geom_tile instead of ggplot .
<code>x</code>	matrix with component intensities in columns
<code>na.rm</code>	see <code>link[base]{min}</code>
<code>legend</code>	should a legend be produced instead of normalized values?
<code>n</code>	of colours to produce in legend
<code>min</code>	numeric with value corresponding to "lowest" colour for each column
<code>max</code>	numeric with value corresponding to "hightest" colour for each column
<code>dx</code>	width of label bar
<code>ny</code>	number of colours in legend
<code>labels</code>	component names
<code>normalize</code>	function to normalize the values.
<code>against</code>	value to mix against (for <code>sub = TRUE</code> only, 1 = white, 0 = black)
<code>sub</code>	subtractive color mixing?

Value

invisible NULL

list with components `ymin`, `max` and `fill` to specify value and fill colour value (still numeric!) for the legend, otherwise the normalized values

ggplot object with legend

character with colours

Author(s)

Claudia Beleites
 Claudia Beleites
 Claudia Beleites

makeraster

makeraster

Description

find an evenly spaced grid for x

Usage

```
makeraster(x, startx, d, newlevels, tol = 0.1)
```

```
fitraster(x, tol = 0.1)
```

Arguments

x	numeric to be fitted with a raster
startx	starting point ("origin") for calculation of the raster
d	step size of the raster
newlevels	levels of the raster
tol	tolerance for rounding to new levels: elements of x within tol of the distance between the levels of the new grid are rounded to the new grid point.

Details

makeraster fits the data to the specified raster.

fitraster tries different raster parameter and returns the raster that covers most of the x values: The differences between the values of x are calculated (possible step sizes). For each of those step sizes, different points are tried (until all points have been covered by a raster) and the parameter combination leading to the best coverage (i.e. most points on the grid) is used.

Note that only differences between the sorted values of x are considered as step size.

Value

list with elements

x	the values of x, possibly rounded to the raster values
levels	the values of the raster

Author(s)

Claudia Beleites

Examples

```

x <- c (sample (1:20, 10), (0 : 5) + 0.5)
raster <- makeraster (x, x [1], 2)
raster
plot (x)
abline (h = raster$levels, col = "#00000040")

## unoccupied levels
missing <- setdiff (raster$levels, raster$x)
abline (h = missing, col = "red")

## points acutally on the raster
onraster <- raster$x %in% raster$levels
points (which (onraster), raster$x [onraster], col = "blue", pch = 20)

raster <- fitraster (x)
raster
plot (x)
abline (h = raster$levels, col = "#00000040")

## unoccupied levels
missing <- setdiff (raster$levels, raster$x)
abline (h = missing, col = "red")

## points acutally on the raster
onraster <- raster$x %in% raster$levels
points (which (onraster), raster$x [onraster], col = "blue", pch = 20)

x <- c (sample (1:20, 10), (0 : 5) + 0.45)
raster <- fitraster (x)
raster
plot (x)
abline (h = raster$levels, col = "#00000040")

## unoccupied levels
missing <- setdiff (raster$levels, raster$x)
abline (h = missing, col = "red")

## points acutally on the raster
onraster <- raster$x %in% raster$levels
points (which (onraster), raster$x [onraster], col = "blue", pch = 20)

```

map.sel.poly

*Interactively select a polygon (grid graphics) and highlight points***Description**

Click the points that should be connected as polygon. Input ends with right click (see [grid.locator](#)). Polygon will be drawn closed.

Usage

```
map.sel.poly(data, pch = 19, size = 0.3, ...)
```

```
sel.poly(pch = 19, size = 0.3, ...)
```

Arguments

data	hyperSpec object for plotting map or list returned by plotmap
pch	symbol to display the points of the polygon for sel.poly
size	size for polygon point symbol for sel.poly
...	further arguments for grid.points and grid.lines

Details

map.sel.poly is a convenience wrapper for [plotmap](#), [sel.poly](#), and [point.in.polygon](#). For customized plotting, the plot can be produced by [plotmap](#), [plotvoronoi](#) or [levelplot](#), and the result of that plot command handed over to map.sel.poly, see the example below.

If even more customized plotting is required, [sel.poly](#) should be used (see example).

Value

map.sel.poly: array of indices for points within the selected polygon

sel.poly: n x 2 matrix with the corner points of the polygon

Author(s)

Claudia Beleites, Sebastian Mellor

Claudia Beleites

See Also

[grid.locator](#), [map.identify](#)

[grid.locator](#)

Examples

```
if (interactive()){
## convenience wrapper
map.sel.poly (chondro)

## customized version
data <- sample (chondro [, , 1004 - 2i ~ 1004 + 2i], 300)

plotdata <- plotvoronoi (data, clusters ~ y * x, col.regions = alois.palette ())
print (plotdata)
map.sel.poly (plotdata)

## even more customization:
```

```

plotvoronoi (data)

## interactively retrieve polygon
polygon <- sel.poly ()

## find data points within polygon
require ("sp")
i.sel <- which (point.in.polygon (data$x, data$y, polygon [, 1], polygon [, 2]) > 0)

## work with selected points
grid.points (unit (data$x [i.sel], "native"), unit (data$y [i.sel], "native"))
}

```

mark.dendrogram

Mark groups in [hclust](#) dendrograms

Description

Groups are marked by colored rectangles as well as by their levels.

Usage

```

mark.dendrogram(
  dendrogram,
  groups,
  col = seq_along(unique(groups)),
  pos.marker = 0,
  height = 0.025 * max(dendrogram$height),
  pos.text = -2.5 * height,
  border = NA,
  text.col = "black",
  label,
  label.right = TRUE,
  ...
)

```

Arguments

dendrogram	the dendrogram
groups	factor giving the the groups to mark
col	vector with colors for each group
pos.marker	top of the marker rectangle
height	height of the marker rectangle
pos.text	position of the text label
border	see text
text.col	color (vector) giving the color for the text markers

label	side label see example
label.right	should the side labels be at the right side?
...	handed to <code>rect</code> and <code>text</code>

Details

The dendrogram should be plotted separately, see the example.

Author(s)

Claudia Beleites

Examples

```
dend <- hclust (pearson.dist (laser[[ ]]))
par (xpd = TRUE, mar = c (5.1, 4, 4, 3)) # allows plotting into the margin
plot (dend, hang = -1, labels = FALSE)

## mark clusters
clusters <- as.factor (cutree (dend, k = 4))
levels (clusters) <- LETTERS [1 : 4]
mark.dendrogram (dend, clusters, label = "cluster")

## mark independent factor
mark.dendrogram (dend, as.factor (laser [,405.36] > 11000),
pos.marker = -0.02, pos.text = - 0.03)

## mark continuous variable: convert it to a factor and omit labels
mark.dendrogram (dend, cut (laser [, 405.36]), 100, alois.palette (100),
pos.marker = -.015, text.col = NA,
label = expression (I [lambda == 405.36~nm]), label.right = FALSE)
```

markpeak

Mark peak Marks location of the first spectrum at the data point closest to the specified position on the current plot.

Description

Mark peak

Marks location of the *first* spectrum at the data point closest to the specified position on the current plot.

Usage

```
markpeak(spc, xpos, col = "red")
```


Arguments

spc	the hyperSpec object
xpos	position of the peak(s) in current x-axis units
col	color of the markers and text

Author(s)

R. Kiselev

Examples

```
plot (chondro [7])
markpeak (chondro [7], 1662)
```

Math2,hyperSpec-method

Math Functions for hyperSpec Objects

Description

Mathematical functions for hyperSpec Objects.

Usage

```
## S4 method for signature 'hyperSpec'
Math2(x, digits)

## S4 method for signature 'hyperSpec'
log(x, base = exp(1), ...)

## S4 method for signature 'hyperSpec'
Math(x)
```

Arguments

x	the hyperSpec object
digits	integer stating the rounding precision
base	base of logarithm
...	ignored

Details

The functions abs, sign, sqrt, floor, ceiling, trunc, round, signif, exp, log, expm1, log1p, cos, sin, tan, acos, asin, atan, cosh, sinh, tanh, acosh, asinh, atanh, lgamma, gamma, digamma, trigamma, cumsum, cumprod, cummax, cummin for hyperSpec objects.

Value

a hyperSpec object

Author(s)

C. Beleites

See Also

[S4groupGeneric](#) for group generic methods.

[Math](#) for the base math functions.

[Arith](#) for arithmetic operators, [Comparison](#) for comparison operators, and [Summary](#) for group generic functions working on hyperSpec objects.

Examples

```
log (flu)
```

matlab.palette	<i>Matlab-like Palettes Two palettes going from blue over green to red, approximately as the standard palette of Matlab does. The second one has darker green values and is better suited for plotting lines on white background.</i>
----------------	---

Description

Matlab-like Palettes Two palettes going from blue over green to red, approximately as the standard palette of Matlab does. The second one has darker green values and is better suited for plotting lines on white background.

Usage

```
matlab.palette(n = 100, ...)
```

```
matlab.dark.palette(n = 100, ...)
```

```
alois.palette(n = 100, ...)
```

Arguments

n	the number of colors to be in the palette.
...	further arguments are handed to rainbow (alois.palette: colorRampPalette)

Value

A vector containing the color values in the form "#rrbbggaa".

Author(s)

C. Beleites and A. Bonifacio

See Also

[rainbow](#)

Examples

```
plotmap (chondro [,, 778], col.regions = matlab.palette ())
```

```
plot (flu, col = matlab.dark.palette (nrow (flu)))
```

```
plotmap (chondro, col = alois.palette)
```

mean_sd,numeric-method

Mean and Standard Deviation Calculate mean and standard deviation, and mean, mean $\pm$ one standard deviation, respectively.

Description

These functions are provided for convenience.

Usage

```
## S4 method for signature 'numeric'
mean_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'matrix'
mean_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'hyperSpec'
mean_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'numeric'
mean_pm_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'matrix'
mean_pm_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'hyperSpec'
mean_pm_sd(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'hyperSpec'
mean(x, na.rm = TRUE, ...)
```

```
## S4 method for signature 'hyperSpec'
quantile(x, probs = seq(0, 1, 0.5), na.rm = TRUE, names = "num", ...)
```

Arguments

<code>x</code>	a numeric vector
<code>na.rm</code>	handed to mean and sd
<code>...</code>	ignored (needed to make function generic)
<code>probs</code>	the quantiles, see quantile
<code>names</code>	"pretty" results in percentages (like quantile 's <code>names = TRUE</code>), "num" results in the row names being <code>as.character(probs)</code> (good for <code>ggplot2</code> getting the order of the quantiles right). Otherwise, no names are assigned.

Value

`mean_sd` returns a vector with two values (mean and standard deviation) of `x`.

`mean_sd(matrix)` returns a matrix with the mean spectrum in the first row and the standard deviation in the 2nd.

`mean_sd` returns a `hyperSpec` object with the mean spectrum in the first row and the standard deviation in the 2nd.

`mean_pm_sd` returns a vector with 3 values: mean - 1 sd, mean, mean + 1 sd

`mean_pm_sd(matrix)` returns a matrix containing mean - sd, mean, and mean + sd rows.

For `hyperSpec` objects, `mean_pm_sd` returns a `hyperSpec` object containing mean - sd, mean, and mean + sd spectra.

For `hyperSpec` object, `mean` returns a `hyperSpec` object containing the mean spectrum.

For `hyperSpec` object, `quantile` returns a `hyperSpec` object containing the respective quantile spectra.

Author(s)

C. Beleites

See Also

[mean](#), [sd](#)

[mean](#), [sd](#)

[quantile](#)

Examples

```
mean_sd (flu [, , 405 ~ 410])
```

```
mean_sd (flu$spc)
```

```
mean_sd (flu)
```

```

mean_pm_sd (flu$c)

mean_pm_sd (flu$spc)

mean_pm_sd (flu)

plot (mean (chondro))

plot (quantile (chondro))

```

```
merge,hyperSpec,hyperSpec-method
```

Merge hyperSpec objects

Description

Merges two hyperSpec objects and [cbinds](#) their spectra matrices, or merges additional extra data into a hyperSpec object.

Usage

```

## S4 method for signature 'hyperSpec,hyperSpec'
merge(x, y, ...)

## S4 method for signature 'hyperSpec,data.frame'
merge(x, y, ...)

## S4 method for signature 'data.frame,hyperSpec'
merge(x, y, ...)

```

Arguments

x	a hyperSpec object or data.frame
y	a hyperSpec object or data.frame (including derived classes like tibble)
...	handed to merge.data.frame

Details

After merging, the spectra matrix can contain duplicates, and is not ordered according to the wavelength.

If the wavelength axis should be ordered, use [orderw1](#).

If a hyperSpec object and a data.frame are merged, the result is of the class of the first (x) object.

Author(s)

C. Beleites

See Also

[merge](#).

[collapse](#) combines hyperSpec objects that do not share the wavelength axis. [rbind](#), and [cbind](#) for combining hyperSpec objects that.

Examples

```
merge (chondro [1:10,, 600], chondro [5:15,, 600], by = c("x", "y"))$.
tmp <- merge (chondro [1:10,, 610], chondro [5:15,, 610],
             by = c("x", "y"), all = TRUE)

tmp$.
wl (tmp)

## remove duplicated wavelengths:
approxfun <- function (y, wl, new.wl){
  approx (wl, y, new.wl, method = "constant",
         ties = function (x) mean (x, na.rm = TRUE)
        )$y
}

merged <- merge (chondro [1:7,, 610 ~ 620], chondro [5:10,, 615 ~ 625], all = TRUE)
merged$.
merged <- apply (merged, 1, approxfun,
                wl = wl (merged), new.wl = unique (wl (merged)),
                new.wavelength = "new.wl")

merged$.

## merging data.frame into hyperSpec object => hyperSpec object
y <- data.frame (filename = sample (flu$filename, 4, replace = TRUE), cpred = 1:4)
y
tmp <- merge (flu, y)
tmp$.

## merging hyperSpec object into data.frame => data.frame
merge (y, flu)
```

ncol,hyperSpec-method *The Number of Rows (Spectra), Columns, and Data Points per Spectrum of an hyperSpec Object*

Description

ncol returns the number of columns in x@data. I.e. the number of columns with additional information to each spectrum (e.g. "x", "y", ...) + 1 (for column spc containing the spectra).

Usage

```
## S4 method for signature 'hyperSpec'
ncol(x)

## S4 method for signature 'hyperSpec'
nrow(x)

nwl(x)

## S4 method for signature 'hyperSpec'
dim(x)

## S4 method for signature 'hyperSpec'
length(x)
```

Arguments

x a hyperSpec object

Value

nrow, ncol, nwl, and length, return an integer.
dim returns a vector of length 3.

Author(s)

C. Beleites

See Also

[ncol](#)
[nrow](#)
[dim](#)
[length](#)

Examples

```
ncol (chondro)
nrow (chondro)

nwl (chondro)
dim (chondro)
length (chondro)
```

normalize01	<i>normalization for mixed colors</i>
-------------	---------------------------------------

Description

Normalize numbers -> [0, 1]

Usage

```
normalize01(x, ...)  
  
## S4 method for signature 'matrix'  
normalize01(x, tolerance = hy.getOption("tolerance"))  
  
## S4 method for signature 'numeric'  
normalize01(x, tolerance = hy.getOption("tolerance"))  
  
## S4 method for signature 'hyperSpec'  
normalize01(x, ...)
```

Arguments

x	vector with values to transform
...	additional parameters such as tolerance handed down.
tolerance	tolerance level for determining what is 0 and 1

Details

The input x is mapped to [0, 1] by subtracting the minimum and subsequently dividing by the maximum. If all elements of x are equal, 1 is returned.

Value

vector with x values mapped to the interval [0, 1]

Author(s)

C. Beleites

See Also

[wl.eval](#), [vanderMonde](#)

orderwl	<i>Sorting the Wavelengths of a hyperSpec Object Rearranges the hyperSpec object so that the wavelength vector is in increasing (or decreasing) order.</i>
---------	--

Description

The wavelength vector is sorted and the columns of the spectra matrix are rearranged accordingly.

Usage

```
orderwl(x, na.last = TRUE, decreasing = FALSE)
```

Arguments

x The hyperSpec object.
na.last, decreasing Handed to [order](#).

Value

A hyperSpec object.

Author(s)

C. Beleites

See Also

[order](#)

Examples

```
## Example 1: different drawing order in plotspc
spc <- new ("hyperSpec", spc = matrix (rnorm (5) + 1:5, ncol = 5))
spc <- cbind (spc, spc+.5)

plot (spc, "spc")
text (wl (spc), spc [[], as.character (1:10))
spc <- orderwl (spc)
plot (spc, "spc")
text (wl (spc), spc [[], as.character (1:10))

## Example 2
spc <- new ("hyperSpec", spc = matrix (rnorm (5)*2 + 1:5, ncol = 5))
spc <- cbind (spc, spc)

plot (seq_len(nwl(spc)), spc[[], type = "b")
spc[[]]
```

```

spc <- orderwl (spc)
lines (seq_len(nwl(spc)), spc[[[]], type = "l", col = "red")
spc[[[]]

```

paracetamol	<i>Paracetamol Spectrum A Raman spectrum of a paracetamol tablet.</i>
-------------	---

Description

Paracetamol Spectrum A Raman spectrum of a paracetamol tablet.

Format

The spectrum was acquired with a Renishaw InVia spectrometer from 100 to 3200 cm⁻¹ in step scan mode. Thus the spectrum has several overlapping wavelength regions.

Author(s)

C. Beleites

Examples

```

paracetamol

plot (paracetamol)
plotspc (paracetamol, c (min ~ 1750, 2800 ~ max), xoffset = 800,
wl.reverse = TRUE)

```

pearson.dist	<i>Distance based on Pearson's R²</i>
--------------	--

Description

The calculated distance is $D^2 = \frac{1-COR(x')}{2}$

Usage

```
pearson.dist(x)
```

Arguments

x a matrix

Details

The distance between the rows of `x` is calculated. The possible values range from 0 (perfectly correlated) over 0.5 (uncorrelated) to 1 (perfectly anti-correlated).

Value

distance matrix (distance object)

Author(s)

C. Beleites

References

S. Theodoridis and K. Koutroumbas: Pattern Recognition, 3rd ed., p. 495

See Also

[as.dist](#)

Examples

```
pearson.dist (flu [[]])  
pearson.dist (flu)
```

plot-methods

Plotting hyperSpec Objects

Description

Plotting hyperSpec objects. The plot method for hyperSpec objects is a switchyard to [plotspec](#), [plotmap](#), and [plotc](#).

Usage

```
## S4 method for signature 'hyperSpec,missing'  
plot(x, y, ...)  
  
## S4 method for signature 'hyperSpec,character'  
plot(x, y, ...)
```

Arguments

<code>x</code>	the hyperSpec object
<code>y</code>	selects what plot should be produced
<code>...</code>	arguments passed to the respective plot function

Details

It also supplies some convenient abbreviations for much used plots.

If `y` is missing, `plot` behaves like `plot (x, y = "spc")`.

Supported values for `y` are:

"spc" calls `plotspc` to produce a spectra plot.

"spcmeansd" plots mean spectrum +/- one standard deviation

"spcprctl5" plots 16th, 50th, and 84th percentile spectre. If the distributions of the intensities at all wavelengths were normal, this would correspond to `"spcmeansd"`. However, this is frequently not the case. Then `"spcprctl5"` gives a better impression of the spectral data set.

"spcprctl5" like `"spcprctl5"`, but additionally the 5th and 95th percentile spectra are plotted.

"map" calls `plotmap` to produce a map plot.

"voronoi" calls `plotvoronoi` to produce a Voronoi plot (tesselated plot, like `"map"` for `hyperSpec` objects with uneven/non-rectangular grid).

"mat" calls `plotmat` to produce a plot of the spectra matrix (not to be confused with `matplot`).

"c" calls `plotc` to produce a calibration (or time series, depth-profile, or the like)

"ts" plots a time series: abbreviation for `plotc` (`x, use.c = "t"`)

"depth" plots a depth profile: abbreviation for `plotc` (`x, use.c = "z"`)

Author(s)

C. Beleites

See Also

`plotspc` for spectra plots (intensity over wavelength),

`plotmap` for plotting maps, i.e. color coded summary value on two (usually spatial) dimensions.

`plotc`

`plot`

Examples

```
plot (flu)
```

```
plot (flu, "c")
```

```
plot (laser, "ts")
```

```
spc <- apply (chondro, 2, quantile, probs = 0.05)
```

```
spc <- sweep (chondro, 2, spc, "-")
```

```
plot (spc, "spcprctl5")
```

```
plot (spc, "spcprctl5")
```

```
plot (spc, "spcmeansd")
```

plotc	<i>Calibration- and Timeseries Plots, Depth-Profiles and the like plotc plots intensities of a hyperSpec object over another dimension such as concentration, time, or a spatial coordinate.</i>
-------	--

Description

If func is not NULL, the summary characteristic is calculated first by applying func with the respective arguments (in func.args) to each of the spectra. If func returns more than one value (for each spectrum), the different values end up as different wavelengths.

Usage

```
plotc(
  object,
  model = spc ~ c,
  groups = NULL,
  func = NULL,
  func.args = list(),
  ...
)
```

Arguments

object	the hyperSpec object
model	the lattice model specifying the plot
groups	grouping variable, e.g. .wavelength if intensities of more than one wavelength should be plotted
func	function to compute a summary value from the spectra to be plotted instead of single intensities
func.args	further arguments to func
...	further arguments to xyplot .

Details

If the wavelength is not used in the model specification nor in groups, nor for specifying subsets, and neither is func given, then only the first wavelength's intensities are plotted and a warning is issued.

The special column names .rownames and .wavelength may be used.

The actual plotting is done by [xyplot](#).

Author(s)

C. Beleites

See Also[xyplot](#)**Examples**

```
## example 1: calibration of fluorescence
plotc (flu) ## gives a warning

plotc (flu, func = mean)
plotc (flu, func = range, groups = .wavelength)

plotc (flu[, , 450], ylab = expression (I ["450 nm"] / a.u.))

calibration <- lm (spc ~ c, data = flu[, , 450]$.)
summary (calibration)
plotc (flu [, , 450], type = c("p", "r"))

conc <- list (c = seq (from = 0.04, to = 0.31, by = 0.01))
ci <- predict (calibration, newdata = conc, interval = "confidence", level = 0.999)

panel.ci <- function (x, y, ...,
                      conc, ci.lwr, ci.upr, ci.col = "#606060") {
  panel.xyplot (x, y, ...)
  panel.lmline (x, y, ...)
  panel.lines (conc, ci.lwr, col = ci.col)
  panel.lines (conc, ci.upr, col = ci.col)
}

plotc (flu [, , 450], panel = panel.ci,
      conc = conc$c, ci.lwr = ci [, 2], ci.upr = ci [, 3])

## example 2: time-trace of laser emission modes
cols <- c ("black", "blue", "#008000", "red")
wl <- i2wl (laser, c(13, 17, 21, 23))

plotspc (laser, axis.args=list (x = list (at = seq (404.5, 405.8, .1))))
for (i in seq_along (wl))
  abline (v = wl[i], col = cols[i], lwd = 2)

plotc (laser [, , wl], spc ~ t, groups = .wavelength, type = "b",
      col = cols)
```

plotmap

Plot a Map and Identify/Select Spectra in the Map [levelplot](#) functions for hyperSpec objects. An image or map of a summary value of each spectrum is plotted. Spectra may be identified by mouse click.

Description

The model can contain the special column name `.wavelength` to specify the wavelength axis.

Usage

```
plotmap(object, model = spc ~ x * y, func = mean, func.args = list(), ...)

## S4 method for signature 'hyperSpec,missing'
levelplot(x, data, ...)

## S4 method for signature 'formula,hyperSpec'
levelplot(
  x,
  data,
  transform.factor = TRUE,
  ...,
  contour = FALSE,
  useRaster = !contour
)

map.identify(
  object,
  model = spc ~ x * y,
  voronoi = FALSE,
  ...,
  tol = 0.02,
  warn = TRUE
)

plotvoronoi(object, model = spc ~ x * y, use.tripack = FALSE, mix = FALSE, ...)
```

Arguments

<code>object, data</code>	the <code>hyperSpec</code> object
<code>model, x</code>	formula specifying the columns of <code>object</code> that are to be displayed by levelplot
<code>func, func.args</code>	Before plotting, <code>plotmap</code> applies function <code>func</code> with the arguments given in the list <code>func.args</code> to each of the spectra. Thus a single summary value is displayed for each of the spectra. This can be suppressed manually by setting <code>func</code> to <code>NULL</code> . It is automatically suppressed if <code>.wavelength</code> appears in the formula.
<code>...</code>	further arguments are passed down the call chain, and finally to levelplot
<code>transform.factor</code>	If the color-coded variable is a factor, should <code>trellis.factor.key</code> be used to compute the color coding and legend?
<code>contour, useRaster</code>	see levelplot

<code>voronoi</code>	Should the plot for identifying spectra by mouse click be produced by <code>plotmap</code> (default) or <code>plotvoronoi</code> ?
<code>tol</code>	tolerance for <code>map.identify</code> as fraction of the viewport (i.e. in "npc" units)
<code>warn</code>	should a warning be issued if no point is within the specified tolerance? See also details.
<code>use.tripack</code>	Whether package <code>tripack</code> should be used for calculating the voronoi polygons. If FALSE, package <code>deldir</code> is used instead. See details.
<code>mix</code>	For Voronoi plots using package <code>tripack</code> , I experienced errors if the data was spatially ordered. Randomly rearranging the rows of the <code>hyperSpec</code> object circumvents this problem.

Details

`plotmap`, `map.identify`, and the `levelplot` methods internally use the same gateway function to [levelplot](#). Thus `transform.factor` can be used with all of them and the panel function defaults to [panel.levelplot.raster](#) for all three. Two special column names, `.rownames` and `.wavelength` may be used.

`levelplot` plots the spectra matrix.

`plotvoronoi` calls `plotmap` with different default settings, namely the panel function defaults to [panel.voronoi](#). [panel.voronoi](#) depends on either of the packages 'tripack' or 'deldir' being installed. For further information, please consult the help page of [panel.voronoi](#). On the [chondro](#) data set, `plotmap` is roughly 5 times faster than `plotvoronoi` using `tripack`, and ca. 15 times faster than `plotvoronoi` using `deldir`. Package `tripack`, however, is free only for non-commercial use. Also, it seems that `tripack` version hang (R running at full CPU power, but not responding nor finishing the calculation) for certain data sets. In this case, `mix = TRUE` may help.

`map.identify` calls `plotmap` and `plotvoronoi`, respectively and waits for (left) mouse clicks on points. Other mouse clicks end the input.

Unlike [panel.identify](#), the indices returned by `map.identify` are in the same order as the points were clicked. Also, multiple clicks on the same point are returned as multiple entries with the same index.

`map.identify` uses option `debuglevel` similar to [spc.identify](#): `debuglevel == 1` will plot the tolerance window if no data point was inside (and additionally labels the point) while `debuglevel == 2` will always plot the tolerance window.

The `map.sel.*` functions offer further interactive selection, see [map.sel.poly](#).

Value

`map.identify` returns a vector of row indices into object of the clicked points.

The other functions return a lattice object.

Author(s)

C. Beleites

See Also

vignette (plotting), vignette (hyperspec)
[plot](#)
[levelplot](#)
[trellis.factor.key](#) for improved color coding of factors
[hyperSpec options spc.identify map.sel.poly](#)
[panel.voronoi](#)

Examples

```
## Not run:
vignette (plotting)
vignette (hyperspec)

## End(Not run)

levelplot (spc ~ y * x, chondro [,1003]) # properly rotated
plotmap (chondro [,1003])

# plot spectra matrix
levelplot (spc ~ .wavelength * t, laser, contour = TRUE, col = "#00000080")
# see also plotmat

plotmap (chondro, clusters ~ x * y)

# Voronoi plots
smpl <- sample (chondro, 300)
plotmap (smpl, clusters ~ x * y)
if (require (deldir))
  plotvoronoi (smpl, clusters ~ x * y,
               use.tripack = FALSE)
```

plotmat

Plot spectra matrix

Description

plots the spectra matrix.

Usage

```
plotmat(
  object,
  y = ".row",
  ylab,
  col = alois.palette(20),
```

```

    ...,
    contour = FALSE
  )

```

Arguments

object	hyperSpec object
y	character giving the name of the extra data column to label the y axis.
ylab	y axis label, defaults to "row" and the label of the extra data column used for the y axis, respectively.
col	see image
...	further parameters for image
contour	should contour be called instead of image ?

Details

If package `plotrix` is available, a color legend is plotted to the right. The right margin is set to at least 5 lines.

Author(s)

Claudia Beleites

See Also

[image](#), [contour](#), [levelplot](#)

Examples

```

plotmat (laser, col = alois.palette (100))

plot (laser, "mat")

plotmat (laser)
plotmat (laser, contour = TRUE, add = TRUE)

## use different y axis labels

plotmat (laser, "t")

plotmat (laser, laser$t / 3600, ylab = "t / h")

```

plotspc	<i>Plotting Spectra Plot the spectra of a hyperSpec object, i.e. intensity over wavelength. Instead of the intensity values of the spectra matrix summary values calculated from these may be used.</i>
---------	---

Description

This is hyperSpec's main plotting function for spectra plots.

Usually, the stacked argument of `plotspc` will do fine, but if you need fine control over the stacking, you may calculate the y offsets yourself.

Usage

```
plotspc(
  object,
  wl.range = TRUE,
  wl.index = FALSE,
  wl.reverse = FALSE,
  spc.nmax = hy.getOption("plot.spc.nmax"),
  func = NULL,
  func.args = list(),
  stacked = NULL,
  stacked.args = list(),
  add = FALSE,
  bty = "l",
  plot.args = list(),
  col = "black",
  lines.args = list(),
  xoffset = 0,
  yoffset = 0,
  nxticks = 10,
  axis.args = list(),
  break.args = list(),
  title.args = list(),
  fill = NULL,
  fill.col = NULL,
  border = NA,
  polygon.args = list(),
  zeroline = list(lty = 2, col = col),
  debuglevel = hy.getOption("debuglevel")
)

stacked.offsets(
  x,
  stacked = TRUE,
  min.zero = FALSE,
```

```

    add.factor = 0.05,
    add.sum = 0,
    .spc = NULL,
    debuglevel = hy.getOption("debuglevel")
)

```

Arguments

<code>object</code>	the hyperSpec object
<code>wl.range</code>	the wavelength range to be plotted. Either a numeric vector or a list of vectors with different wavelength ranges to be plotted separately. The values can be either wavelengths or wavelength indices (according to <code>wl.index</code>).
<code>wl.index</code>	if TRUE, <code>wl.range</code> is considered to give column indices into the spectra matrix. Defaults to specifying wavelength values rather than indices.
<code>wl.reverse</code>	if TRUE, the wavelength axis is plotted backwards.
<code>spc.nmax</code>	maximal number of spectra to be plotted (to avoid accidentally plotting of large numbers of spectra).
<code>func</code>	a function to apply to each wavelength in order to calculate summary spectra such as mean, min, max, etc.
<code>func.args</code>	list with further arguments for <code>func</code>
<code>stacked</code>	if not NULL, a "stacked" plot is produced, see the example. <code>stacked</code> may be TRUE to stack single spectra. A numeric or factor is interpreted as giving the grouping, character is interpreted as the name of the extra data column that holds the groups.
<code>stacked.args</code>	a list with further arguments to stacked.offsets .
<code>add</code>	if TRUE, the output is added to the existing plot
<code>bty</code>	see par
<code>plot.args</code>	list with further arguments to plot
<code>col</code>	see par . <code>col</code> might be a vector giving individual colors for the spectra.
<code>lines.args</code>	list with further arguments to lines . <code>lines.args\$type</code> defaults to "l".
<code>xoffset</code>	vector with abscissa offsets for each of the <code>wl.ranges</code> . If it has one element less than there are <code>wl.ranges</code> , 0 is padded at the beginning. The values are interpreted as the distance along the wavelength axis that the following parts of the spectra are shifted towards the origin. E.g. if <code>wl.range = list(600 ~ 1800, 2800 ~ 3200)</code> , <code>xoffset = 750</code> would result in a reasonable plot. See also the examples.
<code>yoffset</code>	ordinate offset values for the spectra. May be offsets to stack the spectra (stacked.offsets). Either one for all spectra, one per spectrum or one per group in stacked.
<code>nxticks</code>	hint how many tick marks the abscissa should have.
<code>axis.args</code>	list with further arguments for axis . <code>axis.args\$x</code> should contain arguments for plotting the abscissa, <code>axis.args\$y</code> those for the ordinate (again as lists).

break.args	list with arguments for axis.break .
title.args	list with further arguments to title . title.args may contain two lists, \$x, and \$y to set parameters individually for each axis.
fill	if not NULL, the area between the specified spectra is filled with color col. The grouping can be given as factor or numeric, or as a character with the name of the extra data column to use. If a group contains more than 2 spectra, the first and the last are used. If TRUE spectra n and nrow (spc) - n build a group.
fill.col	character vector with fill color. Defaults to brightened colors from col.
border	character vector with border color. You will need to set the line color col to NA in order see the effect.
polygon.args	list with further arguments to polygon which draws the filled areas.
zeroline	NA or a list with arguments abline , used to plot line (s) marking I = 0. NA suppresses plotting of the line. The line is by default turned off if yoffset is not 0.
debuglevel	if > 0, additional debug output is produced, see options for details
x	a hyperSpec object
min.zero	if TRUE, the lesser of zero and the minimum intensity of the spectrum is used as minimum.
add.factor, add.sum	proportion and absolute amount of space that should be added.
.spc	for internal use. If given, the ranges are evaluated on .spc. However, this may change in future.

Details

New plots are created by [plot](#), but the abscissa and ordinate are drawn separately by [axis](#). Also, [title](#) is called explicitly to set up titles and axis labels. This allows fine-grained customization of the plots.

If package plotrix is available, its function [axis.break](#) is used to produce break marks for cut wavelength axes.

Empty levels of the stacking factor are dropped (as no stacking offset can be calculated in that case.)

Value

plotspc invisibly returns a list with

x	the abscissa coordinates of the plotted spectral data points
y	a matrix the ordinate coordinates of the plotted spectral data points
wavelengths	the wavelengths of the plotted spectral data points

This can be used together with [spc.identify](#).

a list containing

offsets	numeric with the yoffset for each group in stacked
groups	numeric with the group number for each spectrum
levels	if stacked is a factor, the levels of the groups

Author(s)

C. Beleites

C. Beleites

See Also

[plot](#), [axis](#), [title](#), [lines](#), [polygon](#), [par](#) for the description of the respective arguments.

[axis.break](#) for cut marks

See [plot](#) for some predefined spectra plots such as mean spectrum +/- one standard deviation and the like.

[identify](#) and [locator](#) about interaction with plots.

[plotspc](#)

Examples

```
plotspc (flu)

## artificial example to show wavelength axis cutting
plotspc (chondro [sample (nrow (chondro), 50)],
        wl.range = list (600 ~ 650, 1000 ~ 1100, 1600 ~ 1700),
        xoffset = c (0, 300, 450))

plotspc (chondro [sample (nrow (chondro), 50)],
        wl.range = list (600 ~ 650, 1000 ~ 1100, 1600 ~ 1700),
        xoffset = c (300, 450))

## some journals publish Raman spectra backwards
plotspc (chondro [sample (nrow (chondro), 50)], wl.reverse = TRUE)

plotspc (laser[(0:4)*20+1,,], stacked = TRUE)

plotspc (laser, func = mean_pm_sd,
        col = c(NA, "red", "black"), lines.args = list (lwd = 2),
        fill = c (1, NA, 1),
        fill.col = "yellow", border = "blue",
        polygon.args = list (lty = 2, lwd = 4),
        title.args = list (xlab = expression (lambda[emission] / nm),
                          y = list(line = 3.4),
                          col.lab = "darkgreen"),
        axis.args = list (x = list (col = "magenta"), y = list (las = 1))
)

mean.pm.sd <- aggregate (chondro, chondro$clusters, mean_pm_sd)
plot (mean.pm.sd, col = matlab.palette (3), fill = ".aggregate", stacked = ".aggregate")
```

```

mean.pm.sd <- aggregate (chondro, chondro$clusters, mean_pm_sd)

offset <- stacked.offsets (mean.pm.sd, ".aggregate")
plot (mean.pm.sd, fill.col = matlab.palette (3), fill = ".aggregate",
      stacked = ".aggregate")

plot (aggregate (chondro, chondro$clusters, mean), yoffset = offset$offsets,
      lines.args = list (lty = 2, lwd = 2), add = TRUE)

barb <- do.call (collapse, barbiturates [1:3])
plot (barb, lines.args = list (type = "h"), stacked = TRUE,
      stacked.args = list (add.factor = .2))

```

qplotc

*Spectra plotting with ggplot2***Description**

Spectra plotting with ggplot2

Usage

```

qplotc(
  object,
  mapping = aes_string(x = "c", y = "spc"),
  ...,
  func = NULL,
  func.args = list(),
  map.pointonly = FALSE
)

```

Arguments

object	hyperSpec object
mapping	see geom_point
...	handed to geom_point
func	function to summarize the wavelengths, if NULL, only the first wavelength is used
func.args	arguments to func
map.pointonly	if TRUE, mapping will be handed to geom_point instead of ggplot .

Details

These functions are still experimental and may change substantially in future.

Value

a [ggplot](#) object

Author(s)

Claudia Beleites

See Also

[plotc](#)

[ggplotgeom_point](#)

Examples

```
qplotc (flu)
qplotc (flu) + geom_smooth (method = "lm")
```

qplotmap

Spectra plotting with ggplot2

Description

Spectra plotting with ggplot2

Usage

```
qplotmap(
  object,
  mapping = aes_string(x = "x", y = "y", fill = "spc"),
  ...,
  func = mean,
  func.args = list(),
  map.tileonly = FALSE
)
```

Arguments

object	hyperSpec object
mapping	see geom_tile
...	handed to geom_tile
func	function to summarize the wavelengths
func.args	arguments to func
map.tileonly	if TRUE, mapping will be handed to geom_tile instead of ggplot .

Details

These functions are still experimental and may change substantially in future.

Note that qplotmap will currently produce the wrong scales if x or y are discrete.

Value

a [ggplot](#) object

Author(s)

Claudia Beleites

See Also

[plotmap](#)

[ggplotgeom_tile](#)

Examples

```
qplotmap (chondro)
qplotmap (chondro) + scale_fill_gradientn (colours = alois.palette ())
```

qplotmixmap

qplotmap with colour mixing for multivariate overlay

Description

map plot with colour overlay.

Usage

```
qplotmixmap(object, ...)
```

Arguments

object	hyperSpec object
...	handed over to qmixmap and qmixmap

Value

invisible list with ggplot2 objects map and legend

Author(s)

Claudia Beleites

See Also[qmixture](#)**Examples**

```
chondro <- chondro - spc.fit.poly.below (chondro)
chondro <- sweep (chondro, 1, apply (chondro, 1, mean), "/")
chondro <- sweep (chondro, 2, apply (chondro, 2, quantile, 0.05), "-")

qplotmixmap (chondro [,c (940, 1002, 1440)],
             purecol = c (colg = "red", Phe = "green", Lipid = "blue"))
```

qplotspc

*Spectra plotting with ggplot2***Description**

Spectra plotting with ggplot2

Usage

```
qplotspc(
  x,
  wl.range = TRUE,
  ...,
  mapping = aes_string(x = ".wavelength", y = "spc", group = ".rownames"),
  spc.nmax = hy.getOption("ggplot.spc.nmax"),
  map.lineonly = FALSE,
  debuglevel = hy.getOption("debuglevel")
)
```

Arguments

x	hyperSpec object
wl.range	wavelength ranges to plot
...	handed to geom_line
mapping	see geom_line
spc.nmax	maximum number of spectra to plot
map.lineonly	if TRUE, mapping will be handed to geom_line instead of ggplot .
debuglevel	if > 0, additional debug output is produced

Details

These functions are still experimental and may change substantially in future.

Value

a [ggplot](#) object

Author(s)

Claudia Beleites

See Also

[plotspc](#)

[ggplotgeom_line](#)

Examples

```
plotspc (chondro)

plotspc (paracetamol, c (2800 ~ max, min ~ 1800)) + scale_x_reverse (breaks = seq (0, 3200, 400))

plotspc (aggregate (chondro, chondro$clusters, mean),
         mapping = aes (x = .wavelength, y = spc, colour = clusters)) +
  facet_grid (clusters ~ .)

plotspc (aggregate (chondro, chondro$clusters, mean_pm_sd),
         mapping = aes (x = .wavelength, y = spc, colour = clusters, group = .rownames)) +
  facet_grid (clusters ~ .)
```

rbind.fill.matrix	<i>Bind matrices by row, and fill missing columns with NA</i>
-------------------	---

Description

The matrices are bound together using their column names or the column indices (in that order of precedence.) Numeric columns may be converted to character beforehand, e.g. using `format`. If a matrix doesn't have colnames, the column number is used (via [make.names](#)(unique = TRUE)).

This is an enhancement to [rbind](#) which adds in columns that are not present in all inputs, accepts a list of data frames, and operates substantially faster

Usage

```
## S3 method for class 'matrix'
rbind.fill(...)

## S3 method for class 'fill'
rbind(...)
```

Arguments

... data frames/matrices to row bind together

Details

Note that this means that a column with name "X1" is merged with the first column of a matrix without name and so on.

Vectors are converted to 1-column matrices prior to rbind.

Matrices of factors are not supported. (They are anyways quite inconvenient.) You may convert them first to either numeric or character matrices. If a character matrix is merged with a numeric, the result will be character.

Row names are ignored.

The return matrix will always have column names.

Value

a matrix

Author(s)

C. Beleites

See Also

`rbind`, `cbind`, `plyr::rbind.fill()`

Examples

```
A <- matrix (1:4, 2)
B <- matrix (6:11, 2)
A
B
hyperSpec:::rbind.fill.matrix (A, B)

colnames (A) <- c (3, 1)
A
hyperSpec:::rbind.fill.matrix (A, B)

hyperSpec:::rbind.fill.matrix (A, 99)

#' rbind.fill(mtcars[c("mpg", "wt")], mtcars[c("wt", "cyl")])
```

`read.asc.PerkinElmer` *File import filter PerkinElmer ASCII spectra*

Description

Imports a single spectrum in PerkinElmer's ASCII format. This function is experimental.

Usage

```
read.asc.PerkinElmer(file = stop("filename or connection needed"), ...)
```

Arguments

file	filename (or connection)
...	further parameters are handed to read.txt.long

Value

hyperSpec object

read.cytomat	<i>Import for Cytospec mat files</i>
--------------	--------------------------------------

Description

These functions allow to import .mat (Matlab V5) files written by Cytospec.

Usage

```
read.cytomat(...)
```

```
read.mat.Cytospec(file, keys2data = FALSE, blocks = TRUE)
```

Arguments

...	read.cytomat for now hands all arguments to read.mat.Cytospec for backwards compatibility.
file	The complete file name (or a connection to) the .mat file.
keys2data	specifies which elements of the Info should be transferred into the extra data
blocks	which blocks should be read? TRUE reads all blocks.

Details

read.cytomat has been renamed to read.mat.Cytospec and is now deprecated. Use read.mat.Cytospec instead.

Value

hyperSpec object if the file contains a single spectra block, otherwise a list with one hyperSpec object for each block.

Note

This function is an ad-hoc implementation and subject to changes.

Author(s)

C. Beleites

See Also

R.matlab::readMat

read.ENVI

Import of ENVI data as hyperSpec object

Description

This function allows ENVI data import as hyperSpec object.

read.ENVI.Nicolet should be a good starting point for writing custom wrappers for read.ENVI that take into account your manufacturer's special entries in the header file.

Usage

```
read.ENVI(
  file = stop("read.ENVI: file name needed"),
  headerfile = NULL,
  header = list(),
  keys.hdr2data = FALSE,
  x = 0:1,
  y = x,
  wavelength = NULL,
  label = list(),
  block.lines.skip = 0,
  block.lines.size = NULL,
  ...,
  pull.header.lines = TRUE
)

read.ENVI.HySpex(
  file = stop("read.ENVI.HySpex: file name needed"),
  headerfile = NULL,
  header = list(),
  keys.hdr2data = NULL,
  ...
)

read.ENVI.Nicolet(
  file = stop("read.ENVI: file name needed"),
  headerfile = NULL,
  header = list(),
  ...,
  x = NA,
  y = NA,
  nicolet.correction = FALSE
)
```

Arguments

file	complete name of the binary file
headerfile	name of the ASCII header file. If NULL, the name of the header file is guessed by looking for a second file with the same basename as file but hdr or HDR suffix.
header	list with header information, see details. Overwrites information extracted from the header file.
keys.hdr2data	determines which fields of the header file should be put into the extra data. Defaults to none. To specify certain entries, give character vectors containing the lowercase names of the header file entries.
x, y	vectors of form c(offset, step size) for the position vectors, see details.
wavelength, label	lists that overwrite the respective information from the ENVI header file. These data is then handed to initialize
block.lines.skip, block.lines.size	BIL and BIP ENVI files may be read in blocks of lines: skip the first block.lines.skip lines, then read a block of block.lines.size lines. If block.lines.NULL, the whole file is read. Blocks are silently truncated at the end of the file (more precisely: to header\$lines).
...	currently unused by read.ENVI, read.ENVI.Nicolet hands those arguments over to read.ENVI
pull.header.lines	(internal) flag whether multi-line header entries grouped by curly braces should be pulled into one line each.
nicolet.correction	see details

Details

ENVI data usually consists of two files, an ASCII header and a binary data file. The header contains all information necessary for correctly reading the binary file.

I experienced missing header files (or rather: header files without any contents) produced by Bruker Opus' ENVI export.

In this case the necessary information can be given as a list in parameter header instead:

header\$	values	meaning
samples	integer	no of columns / spectra in x direction
lines	integer	no of lines / spectra in y direction
bands	integer	no of wavelengths / data points per spectrum
'data type'		format of the binary file
	1	1 byte unsigned integer
	2	2 byte signed integer
	3	4 byte signed integer
	4	4 byte float
	5	8 byte double

	9	16 (2 x 8) byte complex double
	12	2 byte unsigned integer
'header offset'	integer	number of bytes to skip before binary data starts
interleave		directions of the data cube
	"BSQ"	band sequential (indexing: [sample, line, band])
	"BIL"	band interleave by line (indexing: [sample, line, band])
	"BIP"	band interleave by pixel (indexing: [band, line, sample])
'byte order'	0 or "little"	little endian
	1 or "big"	big endian
	"swap"	swap byte order

Some more information that is not provided by the ENVI files may be given:

Wavelength axis and axis labels in the respective parameters. For more information, see [initialize](#).

The spatial information is by default a sequence from 0 to `header$samples - 1` and `header$lines - 1`, respectively. `x` and `y` give offset of the first spectrum and step size.

Thus, the object's `$x` column is: $(0 : \text{header}\$samples - 1) * x[2] + x[1]$. The `$y` column is calculated analogously.

Nicolet uses some more keywords in their header file. They are interpreted as follows:

description	giving the position of the first spectrum
z plot titles	wavelength and intensity axis units, comma separated
pixel size	interpreted as x and y step size (specify <code>x = NA</code> and <code>y = NA</code>)

These parameters can be overwritten by giving a list with the respective elements in parameter header.

The values in header line description seem to be microns while the pixel size seems to be in microns. If `nicolet.correction` is true, the pixel size values (i.e. the step sizes) are multiplied by 1000.

Value

a `hyperSpec` object

Functions

- `read.ENVI.HySpex:`
- `read.ENVI.Nicolet:`

Author(s)

C. Beleites, testing for the Nicolet files C. Dicko

References

This function was adapted from `caTools::read.ENVI()`:

Jarek Tuszynski (2008). `caTools`: Tools: moving window statistics, GIF, Base64, ROC AUC, etc.. R package version 1.9.

See Also

caTools::read.ENVI()
[textio](#)

read.ini

Read INI files

Description

read.ini reads ini files of the form

Usage

```
read.ini(  
  con = stop("Connection con needed."),  
  skip = NULL,  
  encoding = "unknown"  
)
```

Arguments

con	connection or file name
skip	number of lines to skip before first [section] starts
encoding	see readLines

Details

[section] key = value
into a list.

read.ini sanitizes the element names and tries to convert scalars and comma separated numeric vectors to numeric.

Value

a list with one element per section in the .ini file, each containing a list with elements for the key-value-pairs.

Author(s)

C. Beleites

read.jdx

*JCAMP-DX Import for Shimadzu Library Spectra***Description**

this is a first rough import function for JCAMP-DX spectra.

Usage

```
read.jdx(
  filename = stop("filename is needed"),
  encoding = "",
  header = list(),
  keys.hdr2data = FALSE,
  ...,
  NA.symbols = c("NA", "N/A", "N.A."),
  collapse.multi = TRUE,
  wl.tolerance = hy.getOption("wl.tolerance"),
  collapse.equal = TRUE
)
```

Arguments

filename	file name and path of the .jdx file
encoding	encoding of the JCAMP-DX file (used by base::readLines())
header	list with manually set header values
keys.hdr2data	index vector indicating which header entries should be tranfered into the extra data. Usually a character vector of labels (lowercase, without and dashes, blanks, underscores). If TRUE, all header entries are read.
...	further parameters handed to the data import function, e.g.

parameter	meaning	default
xtol	tolerance for checking calculated x values against checkpoints at beginning of line	XFACTOR
ytol	tolerance for checking Y values against MINY and MAXY	YFACTOR

NA.symbols	character vector of text values that should be converted to NA
collapse.multi	should hyperSpec objects from multispectra files be collapsed into one hyperSpec object (if FALSE, a list of hyperSpec objects is returned).
wl.tolerance, collapse.equal	see collapse

Details

So far, AFFN and PAC formats are supported for simple XYDATA, DATA TABLEs and PEAK TABLEs.

NTUPLES / PAGES are not (yet) supported.

DIF, DUF, DIFDUP and SQZ data formats are not (yet) supported.

Value

hyperSpec object

Note

JCAMP-DX support is incomplete and the functions may change without notice. See vignette ("fileio") and the details section.

Author(s)

C. Beleites with contributions by Bryan Hanson

read.spc

Import for Thermo Galactic's spc file format These functions allow to import Thermo Galactic/Grams .spc files.

Description

Import for Thermo Galactic's spc file format These functions allow to import Thermo Galactic/Grams .spc files.

Usage

```
read.spc(
  filename,
  keys.hdr2data = FALSE,
  keys.log2data = FALSE,
  log.txt = TRUE,
  log.bin = FALSE,
  log.disk = FALSE,
  hdr = list(),
  no.object = FALSE
)
```

Arguments

filename	The complete file name of the .spc file.
keys.hdr2data, keys.log2data	character vectors with the names of parameters in the .spc file's log block (log2xxx) or header (hdr2xxx) that should go into the extra data (yyy2data) of the returned hyperSpec object. All header fields specified in the .spc file format specification (see below) are imported and can be referred to by their de-capitalized names.
log.txt	Should the text part of the .spc file's log block be read?
log.bin, log.disk	Should the normal and on-disk binary parts of the .spc file's log block be read? If so, they will be put as raw vectors into the hyperSpec object's log.
hdr	A list with fileheader fields that overwrite the settings of actual file's header. Use with care, and look into the source code for detailed insight on the elements of this list.
no.object	If TRUE, a list with wavelengths, spectra, labels, log and data are returned instead of a hyperSpec object. This parameter will likely be subject to change in future - use with care.

Value

If the file contains multiple spectra with individual wavelength axes, `read.spc` returns a list of `hyperSpec` objects. Otherwise the result is a `hyperSpec` object.

`read.spc.KaiserMap` returns a `hyperSpec` object with data columns x, y, and z containing the stage position as recorded in the .spc files' log.

Note

Only a restricted set of test files was available for development. Particularly, the w-planes feature could not be tested.

If you have .spc files that cannot be read with these function, don't hesitate to contact the package maintainer with your code patch or asking advice.

Author(s)

C. Beleites

References

Source development kit and file format specification of .spc files.

See Also

[textio](#)

Examples

```
## get the sample .spc files from ftirsearch.com (see above)
## Not run:
# single spectrum
spc <- read.spc ("BENZENE.SPC")
plot (spc)

# multi-spectra .spc file with common wavelength axis
spc <- read.spc ('IG_MULTI.SPC')
spc

# multi-spectra .spc file with individual wavelength axes
spc <- read.spc ("BARBITUATES.SPC")
plot (spc [[1]], lines.args = list (type = "h"))

## End(Not run)
```

read.spc.Kaiser	<i>read Kaiser .spc files</i>
-----------------	-------------------------------

Description

Import functions for Kaiser Optical Systems .spc files

Usage

```
read.spc.Kaiser(files, ..., glob = TRUE)

read.spc.KaiserMap(files, keys.log2data = NULL, ...)

read.spc.KaiserLowHigh(
  files = stop("file names needed"),
  type = c("single", "map"),
  ...,
  glob = TRUE
)
```

Arguments

files	If glob = TRUE, filename can contain wildcards. Thus all files matching the name pattern in filename can be specified.
glob	If TRUE the filename is interpreted as a wildcard containing file name pattern and expanded to all matching file names.
keys.log2data, ...	All further arguments are handed over directly to read.spc .
type	what kind of measurement was done? If "map", read.spc.KaiserMap is used instead of read.spc.Kaiser.

Details

read.spc.Kaiser imports sets of .spc files written by Kaiser Optical Systems' Hologram software. It may also serve as an example how to write wrapper functions for read.spc to conveniently import specialized sets of .spc files.

Value

hyperSpec

Examples

```
## for examples, please see `vignette("fileio", package = "hyperSpec")`.
```

read.spe	<i>Import WinSpec SPE file</i>
----------	--------------------------------

Description

Import function for WinSpec SPE files (file version up to 3.0). The calibration data (polynome and calibration data pairs) for x-axis are automatically read and applied to the spectra. Note that the y-calibration data structure is not extracted from the file since it is not saved there by WinSpec and is always empty.

Usage

```
read.spe(  
  filename,  
  xaxis = "file",  
  acc2avg = F,  
  cts_sec = F,  
  keys.hdr2data = c("exposure_sec", "LaserWavelen", "accumulCount", "numFrames",  
    "darkSubtracted")  
)  
  
.read.spe.header(filename)  
  
spe.showcalpoints(filename, xaxis = "file", acc2avg = F, cts_sec = F)
```

Arguments

filename	Name of the SPE file to read data from
xaxis	Units of x-axis, e.g. "file", "px", "nm", "energy", "raman", ... read.spe function automatically checks if the x-calibration data are available and uses them (if possible) to reconstruct the xaxis in the selected units.

acc2avg	whether to divide the actual data set by the number of accumulations, thus transforming <i>accumulated</i> spectra to <i>averaged</i> spectra. WinSpec does not do this automatically, so the spectral intensity is always proportional to the number of accumulations. The flag @data\$averaged is automatically set to TRUE.
cts_sec	whether to divide the actual data set by the exposure time, thus going to count per second unit.
keys.hdr2data	Which metadata from the file header should be saved to the Data slot of a newly created hyperSpec object

Value

hyperSpec object
 hdr list with key=value pairs

Functions

- `.read.spe.header`: Read only header of a WinSpec SPE file (version 2.5)
- `spe.showcalpoints`: Plot the WinSpec SPE file (version 2.5) and show the calibration points stored inside of it (x-axis calibration)

Author(s)

R. Kiselev, C. Beleites

read.txt.Horiba	<i>Import Horiba Labspec exported ASCII files</i>
-----------------	---

Description

Read ASCII (.txt) files exported by Horiba's Labspec software (LabRAM spectrometers)

Usage

```
read.txt.Horiba(
  file,
  cols = c(spc = "I / a.u.", .wavelength = expression(Delta * tilde(nu)/cm^-1)),
  header = TRUE,
  sep = "\t",
  row.names = NULL,
  check.names = FALSE,
  ...
)

read.txt.Horiba.xy(file, ...)

read.txt.Horiba.t(
```

```

    file,
    header = TRUE,
    sep = "\t",
    row.names = NULL,
    check.names = FALSE,
    ...
)

```

Arguments

file connection (file name and path) to the .txt file
 cols, header, sep, row.names, check.names, ...
 further parameters are handed over to [read.txt.wide](#)

Details

read.txt.Horiba.xy reads maps, i.e. .txt files where the first two columns give x and y coordinates.

Value

hyperSpec object

Author(s)

C. Beleites

read.txt.Shimadzu	<i>Reads Shimadzu GCxGC-qMS - Spectra Files (.txt) as exported by Shimadzu Chrome Solution (v. 2.72) Mass Spectrometer: Shimadzu GCMS-QP 2010 Ultra (www.shimadzu.com)</i>
-------------------	--

Description

Reads Shimadzu GCxGC-qMS - Spectra Files (.txt) as exported by Shimadzu Chrome Solution (v. 2.72) Mass Spectrometer: Shimadzu GCMS-QP 2010 Ultra (www.shimadzu.com)

Usage

```
read.txt.Shimadzu(filename, encoding = "", quiet = TRUE)
```

Arguments

filename file name and path of the .txt file
 encoding encoding of the txt file (used by [readLines](#))
 quiet suppress printing of progress

Value

list of spectra tables

Note

This is a first rough import function and the functions may change without notice.

Author(s)

Bjoern Egert

read.txt.wide	<i>Import/export of hyperSpec objects to/from ASCII files A detailed discussion of hyperSpec's file import and export capabilities is given in vignette "fileio".</i>
---------------	---

Description

Besides [save](#) and [load](#), two general ways to import and export data into hyperSpec objects exist.

Firstly, hyperSpec objects can be imported and exported as ASCII files.

Usage

```
read.txt.wide(
  file = stop("file is required"),
  cols = list(spc = "I / a.u.", .wavelength = expression(lambda/nm)),
  sep = "\t",
  row.names = NULL,
  check.names = FALSE,
  ...
)

read.txt.long(
  file = stop("file is required"),
  cols = list(.wavelength = expression(lambda/nm), spc = "I / a.u."),
  header = TRUE,
  ...
)

write.txt.long(
  object,
  file = "",
  order = c(".rownames", ".wavelength"),
  na.last = TRUE,
  decreasing = FALSE,
  quote = FALSE,
```

```

    sep = "\t",
    row.names = FALSE,
    cols = NULL,
    col.names = TRUE,
    col.labels = FALSE,
    append = FALSE,
    ...
)

write.txt.wide(
  object,
  file = "",
  cols = NULL,
  quote = FALSE,
  sep = "\t",
  row.names = FALSE,
  col.names = TRUE,
  header.lines = 1,
  col.labels = if (header.lines == 1) FALSE else TRUE,
  append = FALSE,
  ...
)

```

Arguments

file	filename or connection
cols	the column names specifying the column order. For data import, a list with elements colname = label; for export a character vector with the colnames. Use wavelength to specify the wavelengths.
check.names	handed to read.table . Make sure this is FALSE, if the column names of the spectra are the wavelength values.
...	arguments handed to read.table and write.table , respectively.
header	the file has (shall have) a header line
object	the hyperSpec object
order	which columns should be ordered ? order is used as index vector into a data.frame with columns given by cols.
na.last	handed to order by write.txt.long.
decreasing	logical vector giving the sort order
quote, sep, col.names, row.names	have their usual meaning (see read.table and write.table), but different default values. For file import, row.names should usually be NULL so that the first column becomes a extra data column (as opposed to row names of the extra data).
col.labels	Should the column labels be used rather than the colnames?
append	Should the output be appended to an existing file?
header.lines	Toggle one or two line header (wavelengths in the second header line) for write.txt.wide

Details

Firstly, hyperSpec objects can be imported and exported as ASCII files.

A second option is using the package [R.matlab](#) which provides the functions [readMat](#) and [writeMat](#). hyperSpec comes with a number of pre-defined functions to import manufacturer specific file formats. For details, see vignette ("fileio").

[read.spc](#) imports Thermo Galactic's .spc file format, and ENVI files may be read using [read.ENVI](#).

These functions are very flexible and provide lots of arguments.

If you use them to read or write manufacturer specific ASCII formats, please consider writing a wrapper function and contributing this function to **hyperSpec**. An example is in the "flu" vignette (see vignette ("flu", package = "hyperSpec").

Note that R accepts many packed formats for ASCII files, see [connections](#). For .zip files, see [unzip](#).

For further information, see the examples below, vignette ("fileio") and the documentation of [R.matlab](#).

A second option is using the package [R.matlab](#) which provides the functions [readMat](#) and [writeMat](#). hyperSpec comes with a number of pre-defined functions to import manufacturer specific file formats. For details, see vignette ("file-io").

[read.spc](#) imports Thermo Galactic's .spc file format, and ENVI files may be read using [read.ENVI](#).

These functions are very flexible and provide lots of arguments.

If you use them to read or write manufacturer specific ASCII formats, please consider writing a wrapper function and contributing this function to **hyperSpec**. An example is in the "flu" vignette (see vignette ("flu", package = "hyperSpec").

Note that R accepts many packed formats for ASCII files, see [connections](#). For .zip files, see [unzip](#).

For further information, see the examples below and the documentation of [R.matlab](#).

Author(s)

C. Beleites

See Also

vignette ("fileio") and <http://hyperspec.r-forge.r-project.org/blob/fileio.pdf>, respectively

[read.table](#) and [write.table](#)

[R.matlab](#) for .mat files

[read.ENVI](#) for ENVI data

[read.spc](#) for .spc files

Manufacturer specific file formats: [read.txt.Renishaw](#)

Examples

```
## Not run: vignette ("file-io")

## export & import matlab files
if (require (R.matlab)) {
  # export to matlab file
  writeMat (paste0 (tempdir(), "/test.mat"),
            x = flu[[[]], wavelength = flu@wavelength,
            label = lapply (flu@label, as.character))

  # reading a matlab file
  data <- readMat (paste0 (tempdir(), "/test.mat"))
  print (data)
  mat <- new ("hyperSpec", spc = data$x,
            wavelength = as.numeric(data$wavelength),
            label = data$label[,1])
}

## ascii export & import

write.txt.long (flu,
  file = paste0 (tempdir(), "/flu.txt"),
  cols = c(".wavelength", "spc", "c"),
  order = c("c", ".wavelength"),
  decreasing = c(FALSE, TRUE))

read.txt.long (file = paste0 (tempdir(), "/flu.txt"),
  cols = list (.wavelength = expression (lambda / nm),
  spc = "I / a.u", c = expression ("/" (c, (mg/l))))))

write.txt.wide (flu, file = paste0 (tempdir(), "/flu.txt"),
  cols = c("c", "spc"),
  col.labels = TRUE, header.lines = 2, row.names = TRUE)

write.txt.wide (flu, file = paste0 (tempdir(), "/flu.txt"),
  col.labels = FALSE, row.names = FALSE)

read.txt.wide (file = paste0 (tempdir(), "/flu.txt"),
  # give columns in same order as they are in the file
  cols = list (spc = "I / a.u",
  c = expression ("/" ("c", "mg/l")),
  filename = "filename",
  # plus wavelength label last
  .wavelength = "lambda / nm"),
  header = TRUE)

## End(Not run)
```

rmmvnorm

*Multivariate normal random numbers***Description**

Interface functions to use [rmvnorm](#) for [hyperSpec-class](#) objects.

Usage

```
rmmvnorm(n, mean, sigma)

## S4 method for signature 'numeric,hyperSpec,matrix'
rmmvnorm(n, mean, sigma)

## S4 method for signature 'numeric,hyperSpec,array'
rmmvnorm(n, mean, sigma)

## S4 method for signature 'numeric,matrix,matrix'
rmmvnorm(n, mean, sigma)

## S4 method for signature 'numeric,matrix,array'
rmmvnorm(n, mean, sigma)
```

Arguments

n	vector giving the numer of cases to generate for each group
mean	matrix with mean cases in rows
sigma	common covariance matrix or array (ncol (mean) x ncol (mean) x nrow (mean)) with individual covariance matrices for the groups.

Details

The mvtnorm method for hyperSpec objects supports producing multivariate normal data for groups with different mean but common covariance matrix, see the examples.

See Also

[rmvnorm](#)
[cov](#) and [pooled.cov](#) about calculating covariance of hyperSpec objects.

Examples

```
## multiple groups, common covariance matrix

if (require ("mvtnorm")){
  pcov <- pooled.cov (chondro, chondro$clusters)
  rnd <- rmmvnorm (rep (10, 3), mean = pcov$mean, sigma = pcov$COV)
```

```

    plot (rnd, col = rnd$.group)
  }

```

sample,hyperSpec-method

Random Samples and Permutations Take a sample of the specified size from the elements of x with or without replacement.

Description

Random Samples and Permutations Take a sample of the specified size from the elements of x with or without replacement.

isample returns an vector of indices, sample returns the corresponding hyperSpec object.

Usage

```

## S4 method for signature 'hyperSpec'
sample(x, size, replace = FALSE, prob = NULL)

isample(x, size = nrow(x), replace = FALSE, prob = NULL)

## S4 method for signature 'data.frame'
sample(x, size, replace = FALSE, prob = NULL, drop = FALSE)

## S4 method for signature 'matrix'
sample(x, size, replace = FALSE, prob = NULL, drop = FALSE)

```

Arguments

x	The hyperSpec object, data.frame or matrix to sample from
size	positive integer giving the number of spectra (rows) to choose.
replace	Should sampling be with replacement?
prob	A vector of probability weights for obtaining the elements of the vector being sampled.
drop	see drop : by default, do not drop dimensions of the result

Value

a hyperSpec object, data.frame or matrix with size rows for sample, and an integer vector for isample that is suitable for indexing (into the spectra) of x.

vector with indices suitable for row-indexing x

Author(s)

C. Beleites

See Also[sample](#)**Examples**

```

sample (flu, 3)

plot (flu, col = "darkgray")
plot (sample (flu, 3), col = "red", add = TRUE)

plot (flu, col = "darkgray")
plot (sample (flu, 3, replace = TRUE), col = "#0000FF80", add = TRUE,
      lines.args = list (lwd = 2));

isample (flu, 3)
isample (flu, 3, replace = TRUE)
isample (flu, 8, replace = TRUE)
sample (cars, 2)
sample (matrix (1:24, 6), 2)

```

scale,hyperSpec-method

Center and scale hyperSpec object

Description

link[base]{scale}s the spectra matrix. scale (x, scale = FALSE) centers the data.

Usage

```

## S4 method for signature 'hyperSpec'
scale(x, center = TRUE, scale = TRUE)

```

Arguments

x	the hyperSpec object
center	if TRUE, the data is centered to colMeans (x), FALSE suppresses centering. Alternatively, an object that can be converted to numeric of length nwl (x) by as.matrix (e.g. hyperSpec object containing 1 spectrum) can specify the center spectrum.
scale	if TRUE, the data is scaled to have unit variance at each wavelength, FALSE suppresses scaling. Alternatively, an object that can be converted to numeric of length nwl (x) by as.matrix (e.g. hyperSpec object containing 1 spectrum) can specify the center spectrum.

Details

Package scale provides a fast alternative for base::[scale](#)

Value

the centered & scaled hyperSpec object

Author(s)

C. Beleites

See Also

[scale](#)

package scale.

Examples

```
## mean center & variance scale
tmp <- scale (chondro)
plot (tmp, "spcmeansd")
plot (sample (tmp, 5), add = TRUE, col = 2)

## mean center only
tmp <- scale (chondro, scale = FALSE)
plot (tmp, "spcmeansd")
plot (sample (tmp, 5), add = TRUE, col = 2)

## custom center
tmp <- sweep (chondro, 1, mean, `/~`)
plot (tmp, "spcmeansd")
tmp <- scale (tmp, center = quantile (tmp, .05), scale = FALSE)
```

seq.hyperSpec

Sequence generation along spectra or wavelengths This function generates sequences along the spectra (rows) or wavelengths of hyperSpec objects.

Description

Note that [wl2i](#) generates sequences of indices along the wavelength axis.

Usage

```
## S3 method for class 'hyperSpec'
seq(x, from = 1, to = nrow(x), ..., index = FALSE)
```


Arguments

x	the hyperSpec object
from, to	arguments handed to <code>seq.int</code>
...	arguments for <code>seq</code> , namely by, length.out
index	should a vector with indices be returned rather than a hyperSpec object?

Details

`seq` had to be implemented as S3 method as the generic has only ... arguments (on which no dispatch with differing types is possible).

`seq_along` is not generic, but returns a sequence of the length of the object. As `hyperSpec` provides a Method `length`, it can be used. The result is a sequence of indices for the spectra.

Value

a numeric or hyperSpec object, depending on index.

Author(s)

C. Beleites

See Also

`wl2i` to construct sequences of wavelength indices.

`seq`

Examples

```
seq (flu, index = TRUE)
seq_along (flu)
seq (flu, length.out = 3, index = TRUE) # return value is numeric, not integer!
seq (flu, by = 2, index = TRUE)        # return value is numeric, not integer!

plot (flu, col = "darkgray")
plot (seq (flu, by = 2), add = TRUE, col= "red")
plot (seq (flu, length.out = 2), add = TRUE, col= "blue")
```

spc.bin

Wavelength Binning In order to reduce the spectral resolution and thus gain signal to noise ratio or to reduce the dimensionality of the spectral data set, the spectral resolution can be reduced.

Description

The mean of every by data points in the spectra is calculated.

Usage

```
spc.bin(spc, by = stop("reduction factor needed"), na.rm = TRUE, ...)
```

Arguments

<code>spc</code>	the hyperSpec object
<code>by</code>	reduction factor
<code>na.rm</code>	decides about the treatment of NAs: if FALSE or 0, the binning is done using <code>na.rm = FALSE</code> if TRUE or 1, the binning is done using <code>na.rm = TRUE</code> if 2, the binning is done using <code>na.rm = FALSE</code> , and resulting NAs are corrected with <code>mean(...)</code> , <code>na.rm = TRUE</code> .
<code>...</code>	ignored

Details

Using `na.rm = TRUE` always takes about twice as long as `na.rm = FALSE`.

If the spectra matrix does not contain too many NAs, `na.rm = 2` is faster than `na.rm = TRUE`.

Value

A hyperSpec object with ceiling (`nwl (spc) / by`) data points per spectrum.

Author(s)

C. Beleites

Examples

```
spc <- spc.bin (flu, 5)

plot (flu[1,,425:475])
plot (spc[1,,425:475], add = TRUE, col = "blue")

nwl (flu)
nwl (spc)
```

spc.fit.poly

*Polynomial Baseline Fitting These functions fit polynomial baselines.***Description**

Both functions fit polynomials to be used as baselines. If `apply.to` is `NULL`, a `hyperSpec` object with the polynomial coefficients is returned, otherwise the polynomials are evaluated on the spectral range of `apply.to`.

Usage

```
spc.fit.poly(
  fit.to,
  apply.to = NULL,
  poly.order = 1,
  offset.wl = !(is.null(apply.to))
)

spc.fit.poly.below(
  fit.to,
  apply.to = fit.to,
  poly.order = 1,
  npts.min = max(round(nwl(fit.to) * 0.05), 3 * (poly.order + 1)),
  noise = 0,
  offset.wl = FALSE,
  max.iter = nwl(fit.to),
  stop.on.increase = FALSE,
  debuglevel = hygetOption("debuglevel")
)
```

Arguments

<code>fit.to</code>	<code>hyperSpec</code> object on which the baselines are fitted
<code>apply.to</code>	<code>hyperSpec</code> object on which the baselines are evaluated If <code>NULL</code> , a <code>hyperSpec</code> object containing the polynomial coefficients rather than evaluated baselines is returned.
<code>poly.order</code>	order of the polynomial to be used
<code>offset.wl</code>	should the wavelength range be mapped to $\rightarrow [0, \text{delta wl}]$? This enhances numerical stability.
<code>npts.min</code>	minimal number of points used for fitting the polynomial
<code>noise</code>	noise level to be considered during the fit. It may be given as one value for all the spectra, or for each spectrum separately.
<code>max.iter</code>	stop at the latest after so many iterations.
<code>stop.on.increase</code>	additional stopping rule: stop if the number of support points would increase, regardless whether <code>npts.min</code> was reached or not.

`debuglevel` additional output: 1 shows `npts.min`, 2 plots support points for the final baseline of 1st spectrum, 3 plots support points for 1st spectrum, 4 plots support points for all spectra.

Details

`spc.fit.poly` calculates the least squares fit of order `poly.order` to the *complete* spectra given in `fit.to`. Thus `fit.to` needs to be cut appropriately.

Value

hyperspec object containing the baselines in the spectra matrix, either as polynomial coefficients or as polynomials evaluated on the spectral range of `apply.to`

Author(s)

C. Beleites

See Also

`vignette("baseline", package = "hyperSpec")`
see [options](#) for more on `debuglevel`

Examples

```
## Not run: vignette("baseline", package = "hyperSpec")

spc <- chondro [1 : 10]
baselines <- spc.fit.poly(spc[, , c(625 ~ 640, 1785 ~ 1800)], spc)
plot(spc - baselines)

baselines <- spc.fit.poly.below (spc)
plot (spc - baselines)

spc.fit.poly.below(chondro [1:3], debuglevel = 1)
spc.fit.poly.below(chondro [1:3], debuglevel = 2)
spc.fit.poly.below(chondro [1:3], debuglevel = 3, noise = sqrt (rowMeans (chondro [[1:3]])))
```

<code>spc.identify</code>	<i>Identifying Spectra and Spectral Data Points</i> This function allows to identify the spectrum and the wavelength of a point in a plot produced by plotspc .
---------------------------	---

Description

This function first finds the spectrum with a point closest to the clicked position (see [locator](#)). The distance to the clicked point is evaluated relative to the size of the tolerance window.

Usage

```

spc.identify(
  x,
  y = NULL,
  wavelengths = NULL,
  ispc = NULL,
  tol.wl = diff(range(x))/200,
  tol.spc = diff(range(y))/50,
  point.fn = spc.point.max,
  formatter = spc.label.default,
  ...,
  cex = 0.7,
  adj = c(0, 0.5),
  srt = 90,
  warn = TRUE
)

spc.point.max(wl, spc, wlclic)

spc.point.default(wl, spc, wlclic)

spc.point.min(wl, spc, wlclic)

spc.point.sqr(wl, spc, wlclic, delta = 1L)

spc.label.default(ispc, wl, spc, digits = 3)

spc.label.wlonly(ispc, wl, spc, digits = 3)

```

Arguments

x	either the abscissa coordinates or the list returned by plotspc
y	the ordinate values. Giving y will override any values from x\$y.
wavelengths	the wavelengths for the data points. Giving wavelengths will override any values from x\$wavelengths.
ispc	if a selection of spectra was plotted, their indices can be given in ispc. In this case ispc [i] is returned rather than i.
tol.wl, tol.spc	tolerance in wavelength and spectral intensity to search around the clicked point. See details.
point.fn	function (wl, spc, wlclic) to determine the actual point to label, see details.
formatter	function (i, wl, spc) that produces the labels. If NULL, no labels are displayed.
...	passed to text in order to produce the labels
cex, adj, srt	see par

warn	Should the user be warned if no point is in the considered window? In addition, see the discussion of option <code>debugLevel</code> in the details. If FALSE, the resulting <code>data.frame</code> will have a row of NAs instead.
wl	the wavelength to label
spc	the intensity to label
wlclick	the clicked wavelength
delta	<code>spc.point.sqr</code> fits the parabola in the window <code>wlclick</code> $\pm$ delta points.
digits	how many digits of the wavelength should be displayed?

Details

In a second step, `max.fn` searches for the actual point to label within the specified wavelength window of that spectrum. This allows to label maxima (or minima) without demanding too precise clicks. Currently, the following functions to determine the precise point:

<code>spc.point.default</code>	uses the clicked wavelength together with its spectral intensity
<code>spc.point.max</code>	the point with the highest intensity in the wavelength window
<code>spc.point.min</code>	the point with the lowest intensity in the wavelength window
<code>spc.point.sqr</code>	maximum of a parabola fit through the point with highest intensity and the two surrounding points

`point.fn` is called with the arguments `wl` containing the considered wavelength window, `spc` the respective intensities of the closest spectrum, and `wlclick` the wavelength that was clicked. They return a vector of two elements (wavelength and intensity).

As a last step, a label for the point produced by `formatter` and plotted using `text`. Currently, the following formatters are available:

<code>spc.label.default</code>	spectrum number, wavelength
<code>spc.label.wlonly</code>	wavelength

`formatter` functions receive the number of the spectrum `ispc`, the wavelength `wl`, and the spectral intensity `spc` and produce a character variable suitable for labelling. The predefined formatters surround the label text by spaces in order to easily have an appropriate offset from the point of the spectrum.

The warning issued if no spectral point is inside the tolerance window may be switched off by `warn = FALSE`. In that case, the click will produce a row of NAs in the resulting `data.frame`.

`spc.identify` uses option `debuglevel` to determine whether debugging output should be produced. `debuglevel == 2` will plot the tolerance window for every clicked point, `debuglevel == 1` will plot the tolerance window only if no data point was inside. See [hyperSpec options](#) for details about retrieving and setting options.

You may want to adjust the plot's `ylim` to ensure that the labels are not clipped. As a dirty shortcut, `xpd = NA` may help.

Value

a data.frame with columns

ispc	spectra indices of the identified points, i.e. the rows of the hyperSpec object that was plotted. If ispc is given, ispc [i] is returned rather than i.
wavelengths	the wavelengths of the identified points
spc	the intensities of the identified points

Author(s)

C. Beleites

See Also

[locator](#), [plotspc](#), [hyperSpec options](#)

[map.identify](#) [map.sel.poly](#)

Examples

```
if (interactive ()) {
  ispc <- sample (nrow (laser), 10)
  ispc

  identified <- spc.identify (plotspc (laser[ispc]))

  ## convert to the "real" spectra indices
  ispc [identified$ispc]
  identified$wl
  identified$spc

  ## allow the labels to be plotted into the plot margin
  spc.identify (plotspc (laser[ispc]), ispc = ispc, xpd = NA)

  spc.identify (plotspc (paracetamol, xoffset = 1100,
    wl.range = c (600 ~ 1700, 2900 ~ 3150)),
    formatter = spc.label.wlonly)

  ## looking for minima
  spc.identify (plot (-paracetamol, wl.reverse = TRUE),
    point.fn = spc.point.min, adj = c (1, 0.5))
}
```

spc.loess	<i>loess smoothing interpolation for spectra Spectra can be smoothed and interpolated on a new wavelength axis using loess.</i>
-----------	---

Description

Applying [loess](#) to each of the spectra, an interpolation onto a new wavelength axis is performed. At the same time, the spectra are smoothed in order to increase the signal : noise ratio. See [loess](#) and [loess.control](#) on the parameters that control the amount of smoothing.

Usage

```
spc.loess(spc, newx, enp.target = nwl(spc)/4, surface = "direct", ...)
```

Arguments

spc	the hyperSpec object
newx	wavelength axis to interpolate on
enp.target, surface, ...	parameters for loess and loess.control .

Value

a new hyperspec object.

Author(s)

C. Beleites

See Also

[loess](#), [loess.control](#)

Examples

```
plot (flu, col = "darkgray")
plot (spc.loess(flu, seq (420, 470, 5)), add = TRUE, col = "red")

flu [[3, ]] <- NA_real_
smooth <- spc.loess(flu, seq (420, 470, 5))
smooth [[, ]]
plot (smooth, add = TRUE, col = "blue")
```

spc.NA.approx	<i>Impute missing data points</i>
---------------	-----------------------------------

Description

Replace NAs in the spectra matrix by interpolation. With less than 4 points available linear interpolation of the 2 neighbour points is used. For larger numbers of neighbour points, smoothing interpolation is performed by [smooth.spline](#).

Usage

```
spc.NA.approx(
  spc,
  neighbours = 1,
  w = rep(1, 2 * neighbours),
  df = 1 + .Machine$double.eps,
  spar = NULL,
  debuglevel = hy.getOption("debuglevel")
)

spc.NA.linapprox(...)
```

Arguments

spc	hyperSpec object with spectra matrix containing NAs
neighbours	how many neighbour data points should be used to fit the line
w, df, spar	see smooth.spline
debuglevel	see options
...	ignored

Value

hyperSpec object

Note

The function has been renamed from spc.NA.linapprox to spc.NA.approx

Author(s)

Claudia Beleites

Examples

```
fluNA <- hyperSpec::fluNA
spc.NA.approx (fluNA [, , min ~ 410], debuglevel = 1)
spc.NA.approx (fluNA [1, , min ~ 410], debuglevel = 2)
spc.NA.approx (fluNA [4, , min ~ 410], neighbours = 3, df = 4, debuglevel = 2)
```

`spc.rubberband`*Rubberband baseline correction*

Description

Rubberband baseline

Usage

```
spc.rubberband(spc, ..., upper = FALSE, noise = 0, spline = TRUE)
```

Arguments

<code>spc</code>	hyperSpec object
<code>...</code>	further parameters handed to smooth.spline
<code>upper</code>	logical indicating whether the lower or upper part of the hull should be used
<code>noise</code>	noise level to be taken into account
<code>spline</code>	logical indicating whether the baseline should be an interpolating spline through the support points or piecewise linear.

Details

Baseline with support points determined from a convex hull of the spectrum.

Use `debuglevel >= 1` to obtain debug plots, either directly via function argument or by setting hyperSpec's `debuglevel` option.

Value

hyperSpec object containing the baselines

Note

This function is still experimental

Author(s)

Claudia Beleites

See Also

[spc.fit.poly](#), [spc.fit.poly.below](#)
[vignette\("baseline"\)](#)
[hy.setOptions](#)

Examples

```
plot (paracetamol [,, 175 ~ 1800])
b1 <- spc.rubberband (paracetamol [,, 175 ~ 1800], noise = 300, df = 20)
plot (b1, add = TRUE, col = 2)

plot (paracetamol [,, 175 ~ 1800] - b1)
```

spc.smooth.spline	<i>Spectral smoothing by splines</i>
-------------------	--------------------------------------

Description

Smoothing splines

Usage

```
spc.smooth.spline(spc, newx = wl(spc), ...)
```

Arguments

spc	hyperSpec object
newx	wavelength axis to interpolate on
...	further parameters handed to smooth.spline

Details

Spectral smoothing by splines

Value

hyperSpec object containing smoothed spectra

Note

This function is still experimental

Author(s)

Claudia Beleites

See Also

[spc.loess](#)
[smooth.spline](#)

Examples

```
p <- paracetamol[, , 2200 ~ max]
plot(p, col = "gray")
smooth <- spc.smooth.spline(p[, , c(2200 ~ 2400, 2500 ~ 2825, 3150 ~ max)],
                             wl(paracetamol[, , 2200 ~ max]),
                             df = 4, spar = 1)
plot(smooth, col = "red", add = TRUE)

plot(p - smooth)
```

split	<i>Split a hyperSpec object according to groups split divides the hyperSpec object into a list of hyperSpec objects according to the groups given by f.</i>
-------	---

Description

The hyperSpec objects in the list may be bound together again by `bind("r", list_of_hyperSpec_objects)`.

Usage

```
## S4 method for signature 'hyperSpec'
split(x, f, drop = TRUE)
```

Arguments

x	the hyperSpec object
f	a factor giving the grouping (or a variable that can be converted into a factor by <code>as.factor</code>)
drop	if TRUE, levels off that do not occur are dropped.

Value

A list of hyperSpec objects.

Author(s)

C. Beleites

See Also

[split](#)

Examples

```
dist <- pearson.dist (chondro[[ ]])
dend <- hclust (dist, method = "ward")
z <- cutree (dend, h = 0.15)

clusters <- split (chondro, z)
length (clusters)

# difference in cluster mean spectra
plot (apply (clusters[[2]], 2, mean) - apply (clusters[[1]], 2, mean))
```

subset	<i>subset</i>
--------	---------------

Description

subset for hyperSpec object

Usage

```
## S4 method for signature 'hyperSpec'
subset(x, ...)
```

Arguments

x	hyperSpec object
...	handed to subset (data.frame method)

Value

hyperSpec object containing the respective subset of spectra.

Author(s)

Claudia Beleites

See Also

[subset](#)

Summary

The functions

Description

all, any,

Usage

```
## S4 method for signature 'hyperSpec'  
Summary(x, ..., na.rm = FALSE)
```

```
## S4 method for signature 'hyperSpec'  
is.na(x)
```

```
all_wl(expression, na.rm = FALSE)
```

```
any_wl(expression, na.rm = FALSE)
```

Arguments

x	hyperSpec object
...	further objects
na.rm	logical indicating whether missing values should be removed
expression	expression that evaluates to a logical matrix of the same size as the spectra matrix

Details

sum, prod,

min, max,

range, and

is.na

for hyperSpec objects.

All these functions work on the spectra matrix.

Value

sum, prod, min, max, and range return a numeric, all, any, and is.na a logical.

See Also

[Summary](#) for the base summary functions.

[all.equal](#) and [isTRUE](#)

Examples

```

range (flu)

is.na (flu [, , 405 ~ 410]);

all_wl (flu > 100)

any_wl (flu > 300)
! any_wl (is.na (flu))

```

sweep	<i>Sweep Summary Statistic out of an hyperSpec Object</i> sweep <i>for hyperSpec objects.</i>
-------	---

Description

Calls [sweep](#) for the spectra matrix.

Usage

```

## S4 method for signature 'hyperSpec'
sweep(x, MARGIN, STATS, FUN = "-", check.margin = TRUE, ...)

```

Arguments

x	a hyperSpec object.
MARGIN	direction of the spectra matrix that STATS goes along.
STATS	the summary statistic to sweep out. Either a vector or a hyperSpec object. hyperSpec offers a non-standard convenience function: if STATS is a function, this function is applied first (with the same MARGIN) to compute the statistic. However, no further arguments to the apply function can be given. See the examples.
FUN	the function to do the sweeping, e.g. '-' or '/'.
check.margin	If TRUE (the default), warn if the length or dimensions of STATS do not match the specified dimensions of x. Set to FALSE for a small speed gain when you <i>know</i> that dimensions match.
...	further arguments for FUN

Details

sweep is useful for some spectra preprocessing, like offset correction, subtraction of background spectra, and normalization of the spectra.

Value

A hyperSpec object.

Author(s)

C. Beleites

See Also[sweep](#)**Examples**

```
## Subtract the background / slide / blank spectrum
# the example data does not have spectra of the empty slide,
# so instead the overall composition of the sample is subtracted
background <- apply (chondro, 2, quantile, probs = 0.05)
corrected <- sweep (chondro, 2, background, "-")
plot (corrected, "spcprct15")

## Offset correction
offsets <- apply (chondro, 1, min)
corrected <- sweep (chondro, 1, offsets, "-")
plot (corrected, "spcprct15")

## Min-max normalization (on max amide I)
# the minimum is set to zero by the offset correction.
factor <- apply (corrected, 1, max)
mm.corrected <- sweep (corrected, 1, factor, "/")
plot (mm.corrected, "spcprct15")

## convenience: give function to compute STATS:
mm.corrected2 <- sweep (corrected, 1, max, "/")
plot (mm.corrected2)

## checking
stopifnot (all (mm.corrected2 == mm.corrected))
```

trellis.factor.key	<i>Color coding legend for factors Modifies a list of lattice arguments (as for levelplot, etc.) according to the factor levels. The colorkey will show all levels (including unused), and the drawing colors will be set accordingly.</i>
--------------------	--

Description

trellis.factor.key is used during levelplot-based plotting of factors (for hyperSpec objects) unless transform.factor = FALSE is specified.

Usage

```
trellis.factor.key(f, levelplot.args = list())
```


Arguments

`f` the factor that will be color-coded
`levelplot.args` a list with levelplot arguments

Value

the modified list with levelplot arguments.

Author(s)

C. Beleites

See Also

[levelplot](#)

Examples

```
chondro$z <- factor (rep (c("a", "a", "d", "c"),
                        length.out = nrow (chondro)),
                    levels = letters [1 : 4])

str (trellis.factor.key (chondro$z))

plotmap (chondro, z ~ x * y)

## switch off using trellis.factor.key:
## note that the factor levels are collapsed to c(1, 2, 3) rather than
## c(1, 3, 4)
plotmap (chondro, z ~ x * y, transform.factor = FALSE)

plotmap (chondro, z ~ x * y,
        col.regions = c ("gray", "red", "blue", "dark green"))
```

vanderMonde

Function evaluation on hyperSpec objects

Description

`vandermonde` generates van der Monde matrices, the `hyperSpec` method generates a `hyperSpec` object containing the van der Monde matrix of the wavelengths of a `hyperSpec` object.

Usage

```
vanderMonde(x, order, ...)

## S4 method for signature 'hyperSpec'
vanderMonde(x, order, ..., normalize.wl = normalize01)
```

Arguments

<code>x</code>	object to evaluate the polynomial on
<code>order</code>	of the polynomial
<code>...</code>	hyperSpec method: further arguments to decomposition
<code>normalize.wl</code>	function to transform the wavelengths before evaluating the polynomial (or other function). normalize01 maps the wavelength range to the interval [0, 1]. Use I to turn off.

Details

It is often numerically preferable to map `wl (x)` to [0, 1], see the example.

Value

van der Monde matrix

hyperSpec method: hyperSpec object containing van der Monde matrix as spectra and an additional column `".vdm.order"` giving the order of each spectrum (term).

Author(s)

C. Beleites

See Also

[wl.eval](#) for calculating arbitrary functions of the wavelength,
[normalize01](#)

Examples

```
plot (vanderMonde (flu, 2))
plot (vanderMonde (flu, 2, normalize.wl = I))
```

wc

line/word/character count of ASCII files

Description

`'wc()'` is defunct and will be removed from hyperSpec in future. Consider using `[count_lines()]` instead for line counting.

Usage

```
wc()
```

Author(s)

C. Beleites

See Also`[count_lines()]`

<code>wl</code>	<i>Getting and Setting the Wavelength Axis</i> <code>wl</code> returns the wavelength axis, <code>wl<-</code> sets it.
-----------------	---

Description

The wavelength axis of a `hyperSpec` object can be retrieved and replaced with `wl` and `wl<-`, respectively.

Usage`wl(x)``wl (x, label=NULL, digits=6) <- value`**Arguments**

<code>x</code>	a <code>hyperSpec</code> object
<code>label</code>	The label for the new wavelength axis. See initialize for details.
<code>digits</code>	handed to signif . See details.
<code>value</code>	either a numeric containing the new wavelength vector, or a list with <code>value\$wl</code> containing the new wavelength vector and <code>value\$label</code> holding the corresponding label.

Details

When the wavelength axis is replaced, the `colnames` of `x@data$spc` are replaced by the rounded new wavelengths. `digits` specifies the how many significant digits should be used.

There are two ways to set the label of the new wavelength axis, see the examples. If no label is given, a warning will be issued.

Value

a numeric vector
`hyperSpec` object

Note

wl<- always sets the complete wavelength axis, without changing the columns of the spectra matrix. If you rather want to cut the spectral range, use `[`, for interpolation along the spectral axis see `spc.loess` and for spectral binning `spc.bin`.

Author(s)

C. Beleites

See Also

`signif`
 cutting the spectral range: `[`
 interpolation along the spectral axis: `spc.loess`
 spectral binning: `spc.bin`

Examples

```
wl (laser)

# convert from wavelength to frequency
plot (laser)
wl (laser, "f / Hz") <- 2.998e8 * wl (laser) * 1e9
plot (laser)

# convert from Raman shift to wavelength
# excitation was at 785 nm
plot (chondro [1])
wl (chondro) <- list (wl = 1e7 / (1e7/785 - wl (chondro)), label = expression (lambda / nm))
plot (chondro [1])
```

w1.eval

Evaluate function on wavelengths of hyperSpec object

Description

This is useful for generating certain types of baseline "reference spectra".

Usage

```
wl.eval(x, ..., normalize.wl = I)
```

Arguments

x	hyperSpec object
...	hyperSpec method: expressions to be evaluated
normalize.wl	function to transform the wavelengths before evaluating the polynomial (or other function). Use <code>normalize01</code> to map the wavelength range to the interval [0, 1].

Value

hyperSpec object containing one spectrum for each expression

Author(s)

C. Beleites

See Also

[vanderMonde](#) for polynomials,
[normalize01](#) to normalize the wavenumbers before evaluating the function

Examples

```
plot (wl.eval (laser, exp = function (x) exp (-x)))
```

w12i	<i>Conversion between Wavelength and Spectra Matrix Column Index</i> <i>w12i returns the column indices for the spectra matrix for the given</i> <i>wavelengths. i2wl converts column indices into wavelengths.</i>
------	---

Description

If wavelength is numeric, each of its elements is converted to the respective index. Values outside the range of `x@wavelength` become NA.

Usage

```
w12i(x, wavelength = stop("wavelengths are required."), unlist = TRUE)

i2wl(x, i)
```

Arguments

<code>x</code>	a hyperSpec object
<code>wavelength</code>	the wavelengths to be converted into column indices, either numeric or a formula, see details.
<code>unlist</code>	if multiple wavelength ranges are given, should the indices be unlisted or kept in a list?
<code>i</code>	the column indices into the spectra matrix for which the wavelength is to be computed

Details

If the range is given as a formula (i.e. start ~ end, a sequence index corresponding to start : index corresponding to end is returned. If the wavelengths are not ordered, that may lead to chaos. In this case, call [orderw1](#) first.

Two special variables can be used: min and max, corresponding to the lowest and highest wavelength of x, respectively.

start and end may be complex numbers. The resulting index for a complex x is then

index (Re (x)) + Im (x)

Value

A numeric containing the resulting indices for w12i

i2w1 returns a numeric with the wavelengths

Author(s)

C. Beleites

Examples

```
flu
w12i (flu, 405 : 407)
w12i (flu, 405 ~ 407)

## beginning of the spectrum to 407 nm
w12i (flu, min ~ 407)

## 2 data points from the beginning of the spectrum to 407 nm
w12i (flu, min + 2i ~ 407)

## the first 3 data points
w12i (flu, min ~ min + 2i)

## from 490 nm to end of the spectrum
w12i (flu, 490 ~ max)

## the last 8 data points
w12i (flu, max - 7i ~ max)

## get 450 nm +- 3 data points
w12i (flu, 450 - 3i ~ 450 + 3i)

w12i (flu, 300 : 400) ## all NA:
w12i (flu, 600 ~ 700) ## NULL: completely outside flu's wavelength range

i2w1 (chondro, 17:20)
```

wlconv	<i>Convert different wavelength units</i>
--------	---

Description

The following units can be converted into each other: *nm*, cm^{-1} , *eV*, *THz* and *Raman shift*

Usage

```
wlconv(points, src, dst, laser = NULL)
```

```
nm2raman(x, laser)
```

```
nm2invcm(x, ...)
```

```
nm2ev(x, ...)
```

```
nm2freq(x, ...)
```

```
invcm2raman(x, laser)
```

```
invcm2nm(x, ...)
```

```
invcm2ev(x, ...)
```

```
invcm2freq(x, ...)
```

```
raman2invcm(x, laser)
```

```
raman2nm(x, laser)
```

```
raman2ev(x, laser)
```

```
raman2freq(x, laser)
```

```
ev2raman(x, laser)
```

```
ev2invcm(x, ...)
```

```
ev2nm(x, ...)
```

```
ev2freq(x, ...)
```

```
freq2nm(x, ...)
```

```
freq2invcm(x, ...)
```

```
freq2ev(x, ...)
```

```
freq2raman(x, laser)
```

Arguments

points	data for conversion
src	source unit
dst	destination unit
laser	laser wavelength (required for work with Raman shift)
x	wavelength points for conversion
...	ignored

Functions

- nm2raman: conversion **nanometers** -> **Raman shift (relative wavenumber)**
- nm2invcm: conversion **nanometers** -> **inverse cm (absolute wavenumber)**
- nm2ev: conversion **nanometers** -> **electronvolt**
- nm2freq: conversion **nm** -> **frequency in THz**
- invcm2raman: conversion **inverse cm (absolute wavenumber)** -> **Raman shift (relative wavenumber)**
- invcm2nm: conversion **inverse cm (absolute wavenumber)** -> **nanometers**
- invcm2ev: conversion **inverse cm (absolute wavenumber)** -> **electronvolt**
- invcm2freq: conversion **inverse cm (absolute wavenumber)** -> **frequency in THz**
- raman2invcm: conversion **Raman shift (relative wavenumber)** -> **inverse cm (absolute wavenumber)**
- raman2nm: conversion **Raman shift (relative wavenumber)** -> **nanometers**
- raman2ev: conversion **Raman shift (relative wavenumber)** -> **electronvolt**
- raman2freq: conversion **Raman shift (relative wavenumber)** -> **frequency in THz**
- ev2raman: conversion **electronvolt** -> **Raman shift (relative wavenumber)**
- ev2invcm: conversion **electronvolt** -> **inverse cm (absolute wavenumber)**
- ev2nm: conversion **electronvolt** -> **nanometers**
- ev2freq: conversion **electronvolt** -> **frequency in THz**
- freq2nm: conversion **frequency in THz** -> **nanometers**
- freq2invcm: conversion **frequency in THz** -> **inverse cm (absolute wavenumber)**
- freq2ev: conversion **frequency in THz** -> **electronvolt**
- freq2raman: conversion **frequency in THz** -> **Raman shift (relative wavenumber)**

Author(s)

R. Kiselev

Examples

```
wlconv(3200, "Raman shift", "nm", laser = 785.04)
wlconv( 785, "nm", "invcm")
```

[,hyperSpec-method *Extract and Replace parts of hyperSpec objects*

Description

These Methods allow to extract and replace parts of the hyperSpec object.

Usage

```
## S4 method for signature 'hyperSpec'
x[i, j, l, ..., wl.index = FALSE, drop = FALSE]

## S4 method for signature 'hyperSpec'
x[[i, j, l, ..., wl.index = FALSE, drop = FALSE]]

## S4 method for signature 'hyperSpec'
x$name

## S4 replacement method for signature 'hyperSpec'
x[i, j, ...] <- value

## S4 replacement method for signature 'hyperSpec'
x[[i, j, l, wl.index = FALSE, ...]] <- value

## S4 replacement method for signature 'hyperSpec'
x$name <- value
```

Arguments

x	a hyperSpec Object
i	row index: selects spectra [[and [[<- accept indexing with logical matrix or a n by 2 integer index matrix. In this case the indexing is done inside the spectra matrix. See the examples below.
j	selecting columns of x@data
l	selecting columns of the spectra matrix. If l is numeric, the default behaviour is treating l as wavelengths, <i>not</i> as indices.
...	ignored
wl.index	If TRUE (default), the value(s) in l are treated as column indices for the spectral matrix. Otherwise, the numbers in l are treated as wavelengths and the corresponding column indices are looked up first via wl2i .

drop	For [[: drop unnecessary dimensions, see drop and Extract . Ignored for [, as otherwise invalid hyperSpec objects might result.
name	name of the data column to extract. \$spc yields the spectra matrix.
value	the replacement value

Details

They work with respect to the spectra (rows of `x`), the columns of the data matrix, and the wavelengths (columns of the spectra matrix).

Thus, they can be used for selecting/deleting spectra, cutting the spectral range, and extracting or setting the data belonging to the spectra.

Convenient shortcuts for access of the spectra matrix and the `data.frame` in slot `data` are provided.

Extracting: `[`, `[[`, and `$`.

The version with single square brackets (`[`) returns the resulting `hyperSpec` object.

`[[` yields `data.frame` of slot `@data` of that corresponding `hyperSpec` object returned with the same arguments by `[` if columns were selected (i.e. `j` is given), otherwise the spectra matrix `x@data$spc`.

`$` returns the selected column of the `data.frame` in slot `@data`.

Shortcuts. Three shortcuts to conveniently extract much needed parts of the object are defined:

`x[[]]` returns the spectra matrix.

`x$.` returns the complete slot `@data`, including the spectra matrix in column `$spc`, as a `data.frame`.

`x$. .` returns a `data.frame` like `x$.` but without the spectra matrix.

Replacing: `[<-`, `[[<-`, and `$<-`.

```
## S4 method for signature 'hyperSpec':
```

```
x [i, j, l, \dots] <- value
```

```
## S4 method for signature 'hyperSpec':
```

```
x [[i, j, l, wl.index = FALSE, \dots]] <- value
```

```
## S4 method for signature 'hyperSpec':
```

```
x$name <- value
```

`value` gives the values to be assigned.

For `$`, this can also be a list of the form `list (value = value, label = label)`, with `label` containing the label for data column name.

`[[<-` replaces parts of the spectra matrix.

`[<-` replaces parts of the `data.frame` in slot `x@data`.

`$<-` replaces a column of the `data.frame` in slot `x@data`. The value may be a list with two elements, `value` and `label`. In this case the label of the data column is changed accordingly.

`$. .<-` is again an abbreviation for the `data.frame` without the spectra matrix.

Value

For [, [$<-$], [$<-$], and $\$<-$ a hyperSpec object,
 for [] a matrix or data.frame, and
 for \$ the column of the data.frame @data.
 x[][] returns the complete spectra matrix.
 x\$. returns the complete slot @data,
 x\$. returns the data.frame in @data but without the column @data\$spc containing the spectra matrix.

See Also

[w12i](#) on conversion of wavelength ranges to indices.
[drop](#) and [Extract](#) on drop.

Examples

```
## index into the rows (spectra) -----
## make some "spectra"

## numeric index
plot (flu, "spc", lines.args = list (lty = 2))
plot (flu[1:3], "spc", add = TRUE, col = "red")      # select spectra
plot (flu[-(1:3)], "spc", add = TRUE, col = "blue") # delete spectra

## logic index
plot (flu, "spc", lines.args = list (lty = 2))
index <- rnorm (6) > 0
index
plot (flu[index], "spc", add = TRUE, col = "red")    # select spectra
plot (flu[!index], "spc", add = TRUE, col = "blue") # select spectra

## index into the data columns -----
range (chondro[[], "x"])
colnames (chondro[[], 1])
dim (chondro[[], c(TRUE, FALSE, FALSE)])
chondro$x

## the shortcut functions -----

## extract the spectra matrix
flu[][]

## indexing via logical matrix
summary (flu [[flu < 125]])

## indexing the spectra matrix with index matrix n by 2
ind <- matrix (c (1, 2, 4, 406, 405.5, 409), ncol = 2)
ind
```

```

flu [[ind]]

ind <- matrix (c (1, 2, 4, 4:6), ncol = 2)
ind
flu [[ind, wl.index = TRUE]]

pca <- prcomp (flu[[ ]])

## result is data.frame, if j is given:
result <- flu [, 1:2, 405 ~ 410]
result
class (result)
colnames (result)

## extract the data.frame including the spectra matrix
flu$.
dim(flu$.)
colnames (flu$.)
flu$. $spc

calibration <- lm (spc ~ c, data = flu[, , 450]$.)
calibration

flu$..
colnames (flu$..)

## replacement functions
spc <- flu
spc$.
spc[, "c"] <- 16 : 11
## be careful:
plot (spc)
spc [] <- 6 : 1
spc$..
plot (spc)

spc <- flu [, , 405 ~ 410]
spc [[ ]]
spc [[3]] <- -spc[[3]]
spc [[ ]]
spc [, , 405 : 410] <- -spc[, , 405 : 410]
spc [[ ]]
spc [, , 405 ~ 410] <- -spc[, , 405 ~ 410]

## indexing with logical matrix
spc <- flu [, , min ~ 410]
spc < 125
spc [[spc < 125]] <- NA
spc [[ ]]

## indexing with n by 2 matrix
ind <- matrix (c (1, 2, 4, 406, 405.5, 409), ncol = 2)
ind

```

```
spc [[ind]] <- 3
spc [[]]

ind <- matrix (c (1, 2, 4, 4:6), ncol = 2)
ind
spc [[ind, wl.index = TRUE]] <- 9999
spc [[]]

spc$.
spc$z <- 1 : 6
spc
spc$z <- list (1 : 6, "z / a.u.")
```

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