Package 'hsdar'

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25	package Manage, analyse and simulate hyperspectral data in R	
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Description

The **hsdar** package contains classes and functions to manage, analyse and simulate hyperspectral data. These might be either spectrometer measurements or hyperspectral images through the interface of **raster**.

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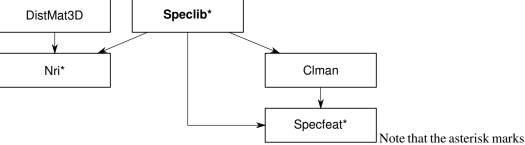
Details

hsdar provides amongst others the following functionality.

• Data handling: **hsdar** is designed to handle even large sets of spectra. Spectra are stored in a Speclib containing, amongst other details, the wavelength and reflectance for each spectrum. **hsdar** further contains functions for plotting spectral data and applying functions to spectra.

- Data manipulation: A variety of established methods for data manipulation such as filter functions (smoothSpeclib), resampling of bands to various satellite sensors (spectralResampling), continuum removal (transformSpeclib), calculations of derivations (derivative.speclib) and extraction of absorption features (cut_specfeat) are implemented.
- Data analysis: Supported methods to analyse vegetation spectra are the calculation of red edge parameters (rededge), vegetation (vegindex) and soil (soilindex) indices as well as ndvilike narrow band indices (nri). hsdar further enables to perform spectral unmixing of spectra (unmix) by use of endmember spectra.
- Data simulation: hsdar has implemented the models PROSAIL 5B (PROSAIL, Jacquemoud et al. 2009) and PROSPECT 5 (PROSPECT, Jacquemoud and Baret 1990) to simulate spectra of canopy and plants.

Several classes are defined and used in **hsdar**. Most of the classes are used and respective objects are created internally. However, the following figure gives an overview which class is used at which stage of processing.



all classes for which wrapper functions for the caret package exist.

To see the preferable citation of the package, type citation("hsdar").

Acknowledgements

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Author(s)

Lukas Lehnert, Hanna Meyer, Joerg Bendix

addcp 5

 - 1	

Manually add fix point to continuum line

Description

This function is used to add an additional fix point to a manually created hull of a single spectrum. This fix point is then used to construct a continuum line.

Usage

```
addcp(x, ispec, cpadd)
```

Arguments

x Object of class Clman.

ispec ID or index of spectrum to be modified.

cpadd Single value or vector of wavelength containing new fix point(s).

Value

Object of class Clman containing the updated version of x.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

```
transformSpeclib, deletecp, getcp, checkhull, makehull, updatecl,
idSpeclib
```

Examples

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)

## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")

## Plot original line
par(mfrow = c(1,2))
plot(spec_clman, ispec = 1, subset = c(2480, 2500))

## Add fix point at 4595 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2495)

## Plot new line</pre>
```

6 apply.DistMat3D

```
plot(spec_clman, ispec = 1, subset = c(2480, 2500))
## Check new hull
hull <- checkhull(spec_clman, 1)
hull$error</pre>
```

apply.DistMat3D

Apply function for class DistMat3D

Description

Apply function to values in a 3-D distance matrix

Usage

```
## S4 method for signature 'DistMat3D'
apply(X, MARGIN, FUN, ...)
```

Arguments

X Object of class 'DistMat3D'.

MARGIN A vector giving the subscripts (dimensions) of the DistMat3D-object which the

function will be applied over (see details).

FUN Function to be applied. Matched with match. fun.

... Further arguments passed to FUN.

Details

The specified function is either applied to the distances of all samples (MARGIN = 1) or to all distances for each sample (MARGIN = 3). In the first case, if X would be replaced by an array of same dimensions the return value would be equal if the following code is applied:

```
apply(X, MARGIN = c(1,2), FUN), where X is an array (see examples).
```

Value

Depending on the length of the return value of the specified function, objects of classes numeric or matrix are returned.

Author(s)

Lukas Lehnert

See Also

```
apply, match.fun, DistMat3D
```

apply.Speclib 7

Examples

```
data(spectral_data)
## Calculate NDVI
ndvi <- nri(spectral_data, b1=800, b2=680)
## Calculate all possible combinations for WorldView-2-8
spec_WV <- spectralResampling(spectral_data, "WorldView2-8",</pre>
                               response_function = FALSE)
nri_WV <- nri(spec_WV, recursive = TRUE)</pre>
class(nri_WV@nri)
## Calculate mean value of all samples for all indices
meanIndexVals <- apply(nri_WV@nri, MARGIN = 1, FUN = mean)</pre>
meanIndexVals
## Same but for array
nri_WV_dat <- as.array(nri_WV@nri)</pre>
meanIndexVals_arr <- apply(nri_WV_dat, MARGIN = c(1, 2), FUN = mean)</pre>
meanSampleVals <- apply(nri_WV@nri, MARGIN = 3, FUN = mean)</pre>
meanSampleVals_arr <- apply(nri_WV_dat, MARGIN = 3, FUN = mean)</pre>
```

apply.Speclib

Apply function for class Speclib

Description

Apply function over all spectra or a subset of spectra

Usage

```
## S4 method for signature 'Speclib'
apply(X, FUN, bySI = NULL, ...)
```

Arguments

Χ	Object of class Speclib
FUN	Function to be applied. Matched with match.fun.
bySI	Character string giving the name of the column in the SI to be used as subsets to apply function FUN on.
	Further arguments passed to FUN.

Value

Object of class Speclib.

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Author(s)

Lukas Lehnert

See Also

```
apply, match.fun, Speclib
```

Examples

```
data(spectral_data)
mean_spectrum <- apply(spectral_data, FUN = mean)
plot(mean_spectrum)

## Same as above but seperately for both seasons
mean_spectra <- apply(spectral_data, FUN = mean, bySI = "season")
plot(mean_spectra, FUN = 1, ylim = c(0,50))
plot(mean_spectra, FUN = 2, new = FALSE)
SI(mean_spectra)</pre>
```

as.hyperSpec

hyperSpec

Description

Conversion from Speclib- to hyperSpec-object

Usage

```
as.hyperSpec(object)
```

Arguments

object

Object of class Speclib.

Value

Object of class hyperSpec.

Note

Package hyperSpec must be installed.

Author(s)

Lukas Lehnert

bandnames 9

See Also

Speclib

bandnames

Handling names of bands

Description

Returning and setting names of bands in Speclib

Usage

```
bandnames(x)
bandnames(x) <- value</pre>
```

Arguments

x Object of class Speclib.

value

Character vector of the same length as nbands(x), or NULL.

Value

For bandnames<-, the updated object. Otherwise a vector giving the name of each band in Speclib is returned.

Author(s)

Lukas Lehnert

See Also

Speclib

Examples

```
data(spectral_data)
bandnames(spectral_data)
```

10 bdri

bdri

Band depth ratio indices

Description

Calculate band depth ratio indices for objects of class Specfeat.

Usage

```
bdri(x, fnumber, index = "ndbi")
```

Arguments

x Object of class Specfeat.

fnumber Integer. Index of feature to modify.

index Method to be applied. Currently, "bdr", "ndbi" and "bna" are available.

Details

Method "bdr" calculates the normalised band depth ratio as

$$bdr = \frac{BD}{Dc},$$

with BD is the band depth calculated by transformSpeclib and Dc is the maximum band depth called band centre. Method "ndbi" calculates the the normalised band depth index as

$$ndbi = \frac{BD - Dc}{BD + Dc}.$$

Method "bna" calculates the band depth normalised to band area as

$$bna = \frac{BD}{Da},$$

where Da is the area of the absorption feature (see feature_properties). For further information see Mutanga and Skidmore (2004).

Value

Object of class specfeat containing the updated version of x.

Author(s)

Lukas Lehnert and Hanna Meyer

References

Mutanga, O. and Skidmore, A. (2004): Hyperspectral band depth analysis for a better estimation of grass biomass (*Cenchrus ciliaris*) measured under controlled laboratory conditions. International Journal of applied Earth Observation and Geoinformation, 5, 87-96

cancer_spectra 11

See Also

transformSpeclib, define.features, specfeat

Examples

cancer_spectra

Hyperspectral samples

Description

Hyperspectral samples from the human larynx

Usage

```
data(cancer_spectra)
```

Format

An object of class Speclib

Details

BIANCA

Author(s)

Bianca Regeling, Lukas Lehnert

caret::createDataPartition-methods

Methods for Function createDataPartition

Description

Methods for function createDataPartition in package **caret**. Please refer to help pages in the **caret**-package for further information.

Methods

```
signature(y = ".CaretHyperspectral") Wrapper method for createDataPartition.

Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
```

caret::createFolds-methods

Methods for Function createFolds and createMultiFolds

Description

Methods for functions createFolds and createMultiFolds in package caret

Methods

```
signature(y = ".CaretHyperspectral") Wrapper methods for createFolds and createMultiFolds.

Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
```

caret::createResample-methods

Methods for Function createResample

Description

Methods for function createResample in package caret

Methods

```
signature(y = ".CaretHyperspectral") Wrapper method for createResample.

Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
```

caret::featurePlot-methods 13

```
caret::featurePlot-methods
```

 ${\it Methods for Function}\ {\it featurePlot}$

Description

Methods for function featurePlot in package caret

Methods

```
signature(x = ".CaretHyperspectral") Wrapper method for featurePlot.
Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
```

caret::gafs

Methods for Function gafs

Description

Methods for function gafs in package **caret**. Please refer to help pages in the **caret**-package for further information.

Usage

```
## S4 method for signature 'Speclib'
gafs(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Nri'
gafs(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Specfeat'
gafs(x, y, cutoff = 0.95, returnData = TRUE, ...)
get_gafs(x)
```

Arguments

х	Object of class Speclib, Nri or Specfeat. For get_gafs, x must be the output of gafs as Speclib or Nri.
У	A numeric or factor vector containing the outcome for each sample. If missing, the response variable set by setResponse is used.
cutoff	The cutoff value of the correlation coefficients between response variables.
returnData	Logical. If TRUE, the updated object of x is returned, otherwise only the result of gafs is returned.
	Further aruments passed to gafs.

Value

If returnData == TRUE, an object of class Speclib or Nri, otherwise an object of class gafs. Note that if x is an object of class Specfeat, the function returns an object of class Speclib containing the relevant transformed band values.

Author(s)

Lukas Lehnert

See Also

gafs

Examples

```
## Not run:
data(spectral_data)

## Set response variable (Chlorophyll content)
spectral_data <- setResponse(spectral_data, "chlorophyll")

## Set additional predictor variables from the SI
spectral_data <- setPredictor(spectral_data, "season")

## Feature selection using genetic algorithms
## Note that this may take some time!
gafs_res <- gafs(spectral_data)

get_gafs(gafs_res)

## End(Not run)</pre>
```

caret::preProcess-methods

Methods for Function preProcess

Description

Methods for function preProcess in package caret. The function is mainly internally required.

Methods

```
signature(x = ".CaretHyperspectral") Wrapper method for preProcess.
Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
```

caret::rfe 15

caret::rfe //	Aethods for Function rfe
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Description

Methods for function rfe in package **caret**. Please refer to help pages in the **caret**-package for further information.

Usage

```
## S4 method for signature 'Speclib'
rfe(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Nri'
rfe(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Specfeat'
rfe(x, y, cutoff = 0.95, returnData = TRUE, ...)
get_rfe(x)
```

Arguments

X	Object of class Speclib, Nri or Specfeat. For get_rfe, x must be the output of rfe as Speclib or Nri.
У	A numeric or factor vector containing the outcome for each sample. If missing, the response variable set by setResponse is used.
cutoff	The cutoff value of the correlation coefficients between response variables.
returnData	Logical. If TRUE, the updated object of x is returned, otherwise only the result of rfe is returned.
	Further aruments passed to rfe.

Value

If returnData == TRUE, an object of class Speclib or Nri, otherwise an object of class rfe. Note that if x is an object of class Specfeat, the function returns an object of class Speclib containing the relevant transformed band values.

Author(s)

Lukas Lehnert

See Also

rfe

16 caret::safs

Examples

```
## Not run:
data(spectral_data)

## Set response variable (Chlorophyll content)
spectral_data <- setResponse(spectral_data, "chlorophyll")

## Set additional predictor variables from the SI
spectral_data <- setPredictor(spectral_data, "season")

## Recursive feature selection
## Note that this may take some time!
rfe_res <- rfe(spectral_data)

get_rfe(rfe_res)

plot(get_rfe(rfe_res))

## End(Not run)</pre>
```

caret::safs

Methods for Function safs

Description

Methods for function safs in package **caret**. Please refer to help pages in the **caret**-package for further information.

Usage

```
## S4 method for signature 'Speclib'
safs(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Nri'
safs(x, y, cutoff = 0.95, returnData = TRUE, ...)
## S4 method for signature 'Specfeat'
safs(x, y, cutoff = 0.95, returnData = TRUE, ...)
get_safs(x)
```

Arguments

X	Object of class Speclib, Nri or Specfeat. For get_safs, x must be the output of safs as Speclib or Nri.
У	A numeric or factor vector containing the outcome for each sample. If missing, the response variable set by setResponse is used.
cutoff	The cutoff value of the correlation coefficients between response variables.

caret::sbf

returnData Logical. If TRUE, the updated object of x is returned, otherwise only the result of safs is returned.

... Further aruments passed to safs.

Value

If returnData == TRUE, an object of class Speclib or Nri, otherwise an object of class safs. Note that if x is an object of class Specfeat, the function returns an object of class Speclib containing the relevant transformed band values.

Author(s)

Lukas Lehnert

See Also

safs

Examples

```
## Not run:
data(spectral_data)

## Set response variable (Chlorophyll content)
spectral_data <- setResponse(spectral_data, "chlorophyll")

## Set additional predictor variables from the SI
spectral_data <- setPredictor(spectral_data, "season")

## Supervised feature selection using simulated annealing
## Note that this may take some time!
safs_res <- safs(spectral_data)

get_safs(safs_res)

plot(get_safs(safs_res))

## End(Not run)</pre>
```

caret::sbf

Methods for Function sbf

Description

Methods for function sbf in package **caret**. Please refer to help pages in the **caret**-package for further information.

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Usage

```
## $4 method for signature 'Speclib'
sbf(x, y, cutoff = 0.95, returnData = TRUE, ...)
## $4 method for signature 'Nri'
sbf(x, y, cutoff = 0.95, returnData = TRUE, ...)
## $4 method for signature 'Specfeat'
sbf(x, y, cutoff = 0.95, returnData = TRUE, ...)
get_sbf(sbf_obj)
```

Arguments

X	Object of class Speclib, Nri or Specfeat.
У	A numeric or factor vector containing the outcome for each sample. If missing, the response variable set by setResponse is used.
cutoff	The cutoff value of the correlation coefficients between response variables.
returnData	Logical. If TRUE, the updated object of x is returned, otherwise only the result of sbf is returned.
	Further aruments passed to sbf.
sbf_obj	Object of class Speclib, Nri or Specfeat as output of sbf-function.

Value

If returnData == TRUE, an object of class Speclib or Nri, otherwise an object of class sbf. Note that if x is an object of class Specfeat, the function returns an object of class Speclib containing the relevant transformed band values.

Author(s)

Lukas Lehnert

See Also

sbf

Examples

```
## Not run:
data(spectral_data)

## Set response variable (Chlorophyll content)
spectral_data <- setResponse(spectral_data, "chlorophyll")

## Set additional predictor variables from the SI
spectral_data <- setPredictor(spectral_data, "season")</pre>
```

caret::setPredictor 19

```
## Selection by filtering
## Note that this may take some time!
sbf_res <- sbf(spectral_data)
get_sbf(sbf_res)
plot(get_sbf(sbf_res))
## End(Not run)</pre>
```

caret::setPredictor

Set predictor variable(s)

Description

Set predictor variable(s) to be used in model-fitting functions of package **caret**. This function can be used to define additional predictor variables stored in the SI of a Speclib- or Nri-object. If the passed object is of class Nri, By default, all Nri-indices (if the passed object is of class Nri) or all bands (if the passed object is of class Speclib) are used as predictors.

Usage

```
## S4 method for signature '.CaretHyperspectral,character'
setPredictor(x, predictor)
```

Arguments

x Object of one of the following classes: Speclib, Nri.

predictor Character vector. Name of additional predictor variable(s) (from the SI).

Value

The updated object.

Author(s)

Lukas Lehnert

See Also

showCaretParameters, setResponse

20 caret::setResponse

Examples

```
## Not run:
data(spectral_data)

## Set "season" as additional predictor variable from the SI
spectral_data <- setPredictor(spectral_data, "season")

## Show caret related parameters stored in the Speclib
showCaretParameters(spectral_data)

## End(Not run)</pre>
```

caret::setResponse Se

Set response variable

Description

Set response variable to be used in model-fitting functions of package **caret**. The response variable must be set upon any model training using a **hsdar**-object in **caret**.

Usage

```
## S4 method for signature '.CaretHyperspectral,character'
setResponse(x, response)
```

Arguments

x Object of one of the following classes: Speclib, Nri.

response Character. Name of response variable (from the SI).

Value

The updated object.

Author(s)

Lukas Lehnert

See Also

showCaretParameters, setPredictor

caret::showCaretParameters 21

Examples

```
## Not run:
data(spectral_data)

## Set response variable (Chlorophyll content)
spectral_data <- setResponse(spectral_data, "chlorophyll")

## Show caret related parameters stored in the Speclib
showCaretParameters(spectral_data)

## End(Not run)</pre>
```

caret::showCaretParameters

Show caret related parameters

Description

Show caret related parameters in objects of classes Speclib, Nri. Several parameters such as predictor and response variables are internally stored and used for model training and validation in the **caret**-package.

Usage

```
showCaretParameters(x)
```

Arguments

Х

Object of one of the following classes: Speclib, Nri.

Author(s)

Lukas Lehnert

See Also

sbf

22 checkhull

Description

Methods for functions train and train. formula in package caret

Methods

```
signature(x = ".CaretHyperspectral") Wrapper method for train.
Note that ".CaretHyperspectral" is a class union containing classes Speclib, Nri.
signature(form = "formula", data = "Speclib") Wrapper method for train.formula to be used with objects of class Speclib.
```

checkhull

Check continuum line

Description

Check if continuum line is intersecting the reflectance curve.

Usage

```
checkhull(x, ispec)
```

Arguments

x Object of class clman.

ispec ID or index of spectrum to be checked.

Value

Object of class list.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

```
transformSpeclib, addcp, deletecp, makehull, updatecl
```

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Examples

```
## Model spectra using PROSAIL
parameter \leftarrow data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)</pre>
## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")</pre>
## Plot original line
par(mfrow = c(1,2))
plot(spec\_clman, ispec = 1, subset = c(2480, 2500))
## Add fix point at 4595 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2495)</pre>
## Plot new line
plot(spec_clman, ispec = 1, subset = c(2480, 2500))
## Check new hull
hull <- checkhull(spec_clman, 1)</pre>
hull$error
## Add fix point at 4596 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2496)</pre>
## Check new hull
hull <- checkhull(spec_clman, 1)</pre>
hull$error
```

Clman

* Clman class

Description

Class to store and handle manual continuum lines.

Details

The class is only required if a continuum line is manually adopted or entirely manually created. This is useful if the automatic approaches are not able to identify absorption features because, for instance, the spectrum has two pronounced maxima within the absortion feature of interest.

Clman is defined as Speclib extended by the following two slots:

- cp: Matrix containing the fix points (continuum points) of each spectrum.
- hull: Matrix containing the hull of each spectrum.

Normally, it is not necessary to manually change the values in any of the slots above. Use the functions addcp and deletecp to change the hulls manually. Functionality for conversion back to a Speclib with the final hull and the transformed spectra provides function updatecl.

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Note

See figure in hsdar-package for an overview of classes in hsdar.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

transformSpeclib, plot, Speclib, addcp, deletecp, updatecl

clman

Methods to create, manipulate and query objects of class 'Clman'.

Description

Methods to create, manipulate and query objects of class 'Clman'. The class 'Clman' is used to store manually defined continuum lines and the associated spectra.

Usage

```
## Creation of objects
## S4 method for signature 'Clman'
initialize(.Object, ...)

## S4 method for signature 'Clman'
spectra(object, ...)

## S4 replacement method for signature 'Clman,data.frame'
spectra(object) <- value

## S4 replacement method for signature 'Clman,matrix'
spectra(object) <- value

## S4 replacement method for signature 'Clman,numeric'
spectra(object) <- value

## S4 method for signature 'Clman'
plot(x, ispec, subset = NULL, numeratepoints = TRUE,
    hull.style = NULL, points.style = list(), ...)</pre>
```

Arguments

.0bject, object Matrix, numeric or array in cases of creation of 'Clman' objects otherwise object of class 'Clman'.

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value	Object of class numeric, matrix or array which is used for replacement of the values in \boldsymbol{x} .
	Arguments passed to createspeclib or plot.default.
x	Object of class clman.
ispec	Name or index of spectrum to be plotted.
subset	Lower and upper spectral limits used for plot.
${\it numerate points}$	Flag if points should be numerated in plot.
hull.style	List of arguments passed to lines to construct the continuum line.
points.style	List of arguments passed to points to construct the continuum points. May be NULL to suppress plotting of fix points.

Value

For spectra<-, the updated object. Otherwise a matrix returning the spectra in the Clman object.

Note

The functions to create objects of class Clman are mainly internally needed by transformSpeclib.

Author(s)

Lukas Lehnert

See Also

```
dist.speclib, Clman, transformSpeclib, plot
```

Examples

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)

## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")

## Return first spectrum
spectra(spec_clman)[1,]

## Plot clman
plot(spec_clman, ispec = 1, subset = c(400, 1000))</pre>
```

26 cor.test

Clman-class

* Clman class

Description

Class to handle continuum removal objects (extends Speclib class).

Details

The class extends Speclibs and adds two additional slots:

- cp: Object of class matrix containing continuum points for all spectra (rows) and bands (columns).
- hull: Object of class matrix containing hull lines for all spectra (rows) and bands (columns).

Note

See figure in hsdar-package for an overview of classes in hsdar.

Author(s)

Lukas Lehnert

See Also

```
Speclib, plot
```

cor.test

Test for association/correlation between nri values and vector of samples

Description

Test for association between paired samples (with one variable being nri-values), using one of Pearson's product moment correlation coefficient, Kendall's tau or Spearman's rho.

Usage

```
## S4 method for signature 'Nri'
cor.test(x, y, ...)
```

cubePlot 27

Arguments

X	Object of class Nri or numerical vector
У	Object of class Nri or numerical vector
	Further arguments passed to cor.test

Details

NRI-values may be used as x and/or as y variable. If x and y are NRI-values the number of samples in both datasets must be equal. For additional information on correlation tests see details in cor.test.

Value

Object of class Nri

Author(s)

Lukas Lehnert

See Also

```
plot, cor.test, glm.nri, lm.nri, getNRI
```

Examples

cubePlot

cubePlot

Description

Plotting 3D cube of hyperspectral data using rgl-package

Usage

28 cut_specfeat

Arguments

X	Object of class HyperSpecRaster.
r	Integer. Index of band used as red channel. If omitted, the band closest to 680 nm is selected.
g	Integer. Index of band used as green channel. If omitted, the band closest to 540 nm is selected.
b	Integer. Index of band used as blue channel. If omitted, the band closest to 470 nm is selected.
ncol	Integer giving the column(s) in x which is/are used to plot the spectral dimension.
nrow	Integer giving the row(s) in x which is/are used to plot the spectral dimension.
sidecol	ColorRamp used to illustrate spectral dimension.
	Further arguments (currently ignored)

Author(s)

Lukas Lehnert

See Also

 ${\bf HyperSpecRaster}$

Examples

```
## Not run:
data(spectral_data)
ras <- HyperSpecRaster(spectral_data, nrow = 9, ncol = 9)
cubePlot(ras)
## End(Not run)</pre>
```

cut_specfeat

Cut absorption features

Description

Function cuts absorption features to a user-specified range.

Usage

```
cut_specfeat(x, ..., fnumber, limits)
```

define.features 29

Arguments

X	An object of class "Specfeat" containing isolated features determined by specfeat.
fnumber	A vector of the positions of the features in x to be cut.
limits	A vector containing the start and end wavelength for each fnumber. The corresponding feature will be cut to this specified range.
	Further arguments passed to generic functions. Currently ignored.

Value

An object of class Specfeat containing the cut features.

Author(s)

Hanna Meyer and Lukas Lehnert

See Also

```
define.features, specfeat, Specfeat
```

Examples

define.features

Definition of absorption features

Description

Function sets the spectral range of absorption features.

30 define.features

Usage

```
define.features(x, tol = 1.0e-7, FWL = NULL)
```

Arguments

X	Object of class Speclib containing the band depth or ratio transformed reflectance spectra.
tol	The tolerance of the band depth which defines a wavelength as a start or end point of a feature. Usually a band depth of 0 or a ratio of 1 indicates feature limits, however, better results are achieved if slightly deviating values are tolerated.
FWL	Optional. If passed, result is directly converted into Specfeat. A vector containing one wavelength per feature to be isolated, e.g. the major absorption features. Features which include these specified wavelengths will be isolated.

Details

Absorption features are defined as the area between local maxima in the reflectance spectra. This function adds the information of the feature limits to the Speclib. Thus, it is a pre-processing step to isolate features.

Value

The updated Speclib containing additional information about the feature limits. If FWL is not NULL, result will be of class Specfeat.

Author(s)

Hanna Meyer and Lukas Lehnert

See Also

```
transformSpeclib, specfeat, Specfeat
```

Examples

deletecp 31

deletecp

Delete fix point

Description

Delete fix point from continuum line.

Usage

```
deletecp(x, ispec, cpdelete)
```

Arguments

x Object of class Clman.

ispec ID or index of spectrum to be modified.

cpdelete Single value or vector of wavelength containing fix point(s) to be deleted.

Value

Object of class Clman containing the updated version of x.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

transformSpeclib, addcp, getcp, checkhull, makehull, updatecl

Examples

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)
## Mask parts not necessary for the example
mask(spec) <- c(1600, 2600)

## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")

## Plot original line
par(mfrow = c(1,2))
plot(spec_clman, ispec = 1, subset = c(1100, 1300))

## Find wavelength of fix point to be deleted
getcp(spec_clman, 1, subset = c(1100, 1300))

## Delete all fix points between 1200 and 1240 nm
spec_clman <- deletecp(spec_clman, 1, c(1200:1240))</pre>
```

32 derivative.speclib

```
## Plot new line
plot(spec_clman, ispec = 1, subset = c(1100, 1300))
## Check new hull
hull <- checkhull(spec_clman, 1)
hull$error</pre>
```

derivative.speclib

Derivation

Description

Calculate derivations of spectra

Usage

```
derivative.speclib(x, m = 1, method = "sgolay", ...)
```

Arguments

x Object of class Speclib.

m Return the m-th derivative of the spectra.

method Character string giving the method to be used. Valid options are "finApprox"

or "sgolay".

... Further arguments passed to sgolayfilt.

Details

Two different methods are available:

• Finite approximation (finApprox):

$$\frac{dr}{d\lambda} = \frac{r(\lambda_i) - r(\lambda_{i+1})}{\Delta\lambda},$$

where r_i is the reflection in band i and $\Delta \lambda$ the spectral difference between adjacent bands.

• Savitzky-Golay derivative computation (sgolay)

Value

Object of class Speclib.

Author(s)

Lukas Lehnert

dim.speclib 33

References

Tsai, F. & Philpot, W. (1998): Derivative analysis of hyperspectral data. Remote Sensing of Environment 66/1. 41-51.

See Also

```
sgolayfilt, vegindex
```

Examples

```
data(spectral_data)
## Calculate 1st derivation
d1 <- derivative.speclib(spectral_data)
## Calculate 2nd derivation
d2 <- derivative.speclib(spectral_data, m = 2)
## Calculate 3rd derivation
d3 <- derivative.speclib(spectral_data, m = 3)
par(mfrow=c(2,2))
plot(spectral_data)
plot(d1)
plot(d2)
plot(d3)</pre>
```

dim.speclib

Dimensions of Speclib

Description

```
Get dimension(s) of Speclib
```

Usage

```
## S4 method for signature 'Speclib'
dim(x)

nspectra(x)
nbands(x)
```

Arguments

Х

Object of class Speclib.

Value

Vector of length = 2 or single integer value.

34 dist.speclib

Author(s)

Lukas Lehnert

See Also

Speclib

Examples

```
data(spectral_data)
dim(spectral_data)
```

dist.speclib

Distance between spectra

Description

Calculation of distance matrices by using one of the various distance measure to compute the distances between the spectra in Speclib. Spectral Angle Mapper (SAM) is calculated with sam giving reference spectra or with sam_distance taking all combinations between spectra in single Speclib into account.

Usage

```
dist.speclib(x, method = "sam", ...)
## Direct call to Spectral Angle Mapper function
sam(x, ref)
sam_distance(x)
```

Arguments

x Object of class Speclib.
 method The distance measure to be used. This must be one of "sam", "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski".
 ref Object of class Speclib containing reference spectra.
 ... Further arguments, passed to other methods.

Details

Available distance measures are "spectral angle mapper" (sam) and all distance measures available in dist. Spectral angle mapper is calculated with the following formula:

$$sam = \cos^{-1}\left(\frac{\sum_{i=1}^{nb} t_i r_i}{\sqrt{\sum_{i=1}^{nb} t_i^2} \sqrt{\sum_{i=1}^{nb} r_i^2}}\right)$$

distMat3D 35

nb is the number of bands in Speclib. t_i and r_i are the reflectances of target and reference spectrum in band i, respectively.

Value

The dist-method for Speclibs returns an object of class "dist". See dist for further information on class "dist". Both other functions return an object of class matrix.

Author(s)

Lukas Lehnert

References

Kruse, F. A.; Lefkoff, A. B.; Boardman, J. W.; Heidebrecht, K. B.; Shapiro, A. T.; Barloon, P. J. & Goetz, A. F. H. (1993). The spectral image processing system (SIPS) – interactive visualization and analysis of imaging spectrometer data. Remote Sensing of Environment, 44, 145-163.

See Also

```
dist, Speclib
```

Examples

distMat3D

Methods to create, manipulate and query objects of class 'Dist-Mat3D'.

Description

Methods to create, manipulate and query objects of class 'DistMat3D'. The following relational operators are defined to compare values between 'DistMat3D'-object(s): <, <=, ==, >, >=

36 distMat3D

Usage

```
## Creation of objects
## S4 method for signature 'numeric'
distMat3D(x, ncol, nlyr)
## S4 method for signature 'matrix'
distMat3D(x, lower_tri = TRUE)
## S4 method for signature 'array'
distMat3D(x, lower_tri = TRUE)
## Conversion methods
## S4 method for signature 'DistMat3D'
as.array(x)
## S4 method for signature 'DistMat3D'
as.matrix(x, lyr = 1)
## Query of properties
## S4 method for signature 'DistMat3D'
dim(x)
## S4 method for signature 'DistMat3D'
ncol(x)
## S4 method for signature 'DistMat3D'
nrow(x)
## Manipulate and query data in objects
## S4 method for signature 'DistMat3D, ANY, ANY'
x[i, j, n]
## S4 replacement method for signature 'DistMat3D, ANY, ANY'
x[i, j, n] \leftarrow value
## S4 method for signature 'DistMat3D'
show(object)
```

Arguments

x,object	Matrix, numeric or array in cases of creation of 'DistMat3D' objects otherwise object of class 'DistMat3D'.
ncol	Number of columns in the new 'DistMat3D' object.
nlyr	Number of layer in the new 'DistMat3D' object.

lower_tri Flag if only the lower triangle is used.

lyr Layer in the 'DistMat3D' object to be transformed into matrix.

DistMat3D-class 37

value Object of class numeric, matrix or array which is used for replacement of the values in x.

i,j,n Subscripts to access data.

Author(s)

Lukas Lehnert

See Also

```
DistMat3D, apply, Nri
```

Examples

```
data(spectral_data)
## Mask channel crossing part (arround 1050 nm) and strong
## water absorption part (above 1350 nm)
mask(spectral_data) <- c(1045, 1055, 1350, 1706)

## Calculate SAM distances (object of class 'dist')
sam_dist <- dist.speclib(subset(spectral_data, season == "summer"))

## Convert to class 'distMat3D'
sam_dist <- distMat3D(as.matrix(sam_dist))
sam_dist</pre>
```

DistMat3D-class

* DistMat3D class

Description

Class to store effectively (large) distance matrices (up to 3D).

Details

Object with 3 slots:

- values: Numerical vector containing distance values
- ncol: Number of columns in the 3D-matrix. Number of columns equals always the number of rows
- nlyr: Number of layers in the 3D-matrix

Note

See figure in hsdar-package for an overview of classes in hsdar.

38 feature_properties

Author(s)

Lukas Lehnert

See Also

distMat3D

feature_properties

Calculation of properties of features

Description

Function to calculate feature properties such as the area, the position of the maximum and several other parameters. This function can only be used for spectral data transformed using any kind of continuum removal (see transformSpeclib).

Usage

feature_properties(x)

Arguments

Х

Object of class Specfeat

Details

The function calculates several parameters:

• area: The feature area is calculated by

$$area_{F_i} = \sum_{k=min(\lambda)}^{max(\lambda)} BD\lambda,$$

with $area_{F_i}$ is the area of the feature i, $min(\lambda)$ is the minimum wavelength of the spectrum, $max(\lambda)$ is the maximum wavelength of the spectrum and BD is the band depth.

- max: Wavelength position of the maximum value observed in the feature.
- Parameters based on half-max values:
 - lo and up: Wavelength position of the lower and upper half-max value.
 - width: Difference between wavelength positions of upper and lower half-max values.
 - gauss_lo: Similarity of the Gauss distribution function and the feature values between the lower half-max and the maximum position. As similarity measurement, the root mean square error is calculated.
 - gauss_up: Same as above but for feature values between the maximum position and the upper half-max.

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Value

An object of class Specfeat containing the properties as (part of the) SI table.

Author(s)

Hanna Meyer \& Lukas Lehnert

See Also

```
define.features, specfeat
```

Examples

get.gaussian.response Gaussian response function

Description

Simulate Gaussian response function for band(s) of a (satellite) sensor. Each band is either defined by center and full-width-half-maximum values or by passing its upper and lower border.

Usage

```
get.gaussian.response(fwhm)
```

Arguments

fwhm

Object of class data. frame with three columns. See details and examples sections.

Details

The characteristics of the sensor must be passed as a data.frame with three columns: first column is used as name for bands, second with lower bounds of channels and third column with upper bounds (5% sensitivity). Alternatively, the data.frame may encompass band centre wavelength and full-width-half-maximum values of the sensor. Function will check the kind of data passed by partially matching the names of the data frame: If any column is named "fwhm" or "center", it is assumed that data are band centre and full-width-half-maximum values.

Value

Data frame with response values for all bands covering the entire spectral range of sensor passed to the function.

Author(s)

Lukas Lehnert

See Also

```
get.sensor.characteristics, spectralResampling
```

```
par(mfrow=c(1,2))
## Plot response function of RapidEye
plot(c(0,1)\sim c(330,1200), type = "n", xlab = "Wavelength [nm]",
     ylab = "Spectral response")
data_RE <- get.gaussian.response(get.sensor.characteristics("RapidEye"))</pre>
xwl_response <- seq.int(attr(data_RE, "minwl"),</pre>
                        attr(data_RE, "maxwl"),
                         attr(data_RE, "stepsize"))
for (i in 1:ncol(data_RE))
 lines(xwl_response, data_RE[,i], col = i)
## Plot original response function
data_RE <- get.sensor.characteristics("RapidEye", TRUE)</pre>
plot(c(0,1)^c(330,1200), type = "n", xlab = "Wavelength [nm]",
     ylab = "Spectral response")
xwl_response <- seq.int(attr(data_RE$response, "minwl"),</pre>
                         attr(data_RE$response, "maxwl"),
                        attr(data_RE$response, "stepsize"))
for (i in 1:nrow(data_RE$characteristics))
 lines(xwl_response, data_RE$response[,i], col = i)
## Simulate gaussian response for arbitrary sensor with 3 bands
sensor <- data.frame(Name = paste("Band_", c(1:3), sep = ""),</pre>
                     center = c(450, 570, 680),
                     fwhm = c(30, 40, 30))
## Plot response function
```

get.sensor.characteristics 41

get.sensor.characteristics

Sensor characteristics

Description

Get channel wavelength of satellite sensor

Usage

```
get.sensor.characteristics(sensor, response_function = FALSE)
```

Arguments

sensor Character or integer. Name or numerical abbreviation of sensor. See 'sensor="help"' or 'sensor=0' for an overview of available sensors.

response_function

If TRUE, the spectral response function is returned

Author(s)

Lukas Lehnert

See Also

```
spectralResampling
```

42 get.sensor.name

get.sensor.name

Satellite sensor name

Description

Get satellite sensor name by integer value

Usage

```
get.sensor.name(sensor)
```

Arguments

sensor

Integer value to match against predefined satellite sensors.

Details

See get.sensor.characteristics to get overview on available satellite sensors.

Value

Name of satellite sensor as character string.

Author(s)

Lukas Lehnert

See Also

```
get.sensor.characteristics
```

```
get.sensor.name(1)
```

getcp 43

getcp

Get fix points

Description

Get fix points of continuum line within spectral range.

Usage

```
getcp(x, ispec, subset = NULL)
```

Arguments

x Object of class Clman.

ispec ID or index of spectrum to be analysed.

subset Vector of length = 2 giving the lower and upper limit of spectral range.

Value

Object of class list containing two elements:

- ptscon: Data frame with wavelength and reflectance of fix points
- ispec: Index of analysed spectrum within passed Clman-object.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

```
transformSpeclib, deletecp, addcp, Clman
```

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)

## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")

## Fix points
spec_cp <- getcp(spec_clman, 1, c(400, 800))
spec_cp</pre>
```

44 getNRI

getNRI

Return nri-values

Description

Return normalized ratio index values giving the wavelength

Usage

```
getNRI(nri, wavelength)
```

Arguments

nri

Object of class 'Nri'

wavelength

Wavelength values where nri is returned. See details section.

Details

Wavelength can be passed in three ways. As the result of nri_best_performance, as a data frame with two columns or as a vector of length 2. In the first two cases, the result will be a data frame (if data frames contain more than one row) with the nri-values of each pair of wavelengths. In the latter case it will be a vector.

Author(s)

Lukas Lehnert

See Also

```
nri, Nri
```

get_reflectance 45

get_refl	ectance	Get reflectance values	

Description

Returns weighted or unweighted reflectance values at wavelength position.

Usage

```
get_reflectance(spectra, wavelength, position, weighted = FALSE, ...)
```

Arguments

spectra	Object of class Speclib or data.frame with reflectance values.	
wavelength	Vector with wavelength values. May be missing if spectra is object of class Speclib.	
position	Numeric value passing the position of reflectance values to be returned in dimensions of the wavelength values.	
weighted	Logical indicating if reflectance values should be interpolated to fit wavelength position. If FALSE the reflectance values of nearest neighbour to passed position are returned.	
	Arguments to be passed to specific functions. Currently ignored.	

Value

A vector with reflectance values for each spectrum is returned. If position falls outside of spectral range of input values, NA values are returned.

Author(s)

Lukas Lehnert \& Hanna Meyer

See Also

```
spectra
```

```
data(spectral_data)
## Simulate multispectral sensor encompassing two bands
## to show effect of weighted and unweighted modes
spectral_data_res <- spectralResampling(spectral_data,
    sensor = data.frame(lb = c(400, 600), ub = c(500, 700)))
## Compare reflectance at 520 nm (in between both bands to</pre>
```

46 glm.nri

glm.nri

(Generalised) Linear models from normalised ratio indices

Description

Build (generalised) linear models of normalised ratio indices as response and predictor variables

Usage

```
lm.nri(formula, preddata = NULL, ...)
glm.nri(formula, preddata = NULL, ...)
```

Arguments

formula Formula for (generalized) linear model

preddata Data frame or speclib containing predictor variables

... Further arguments passed to lm, glm and generic print.default

Details

NRI-values may be used as predictor or response variable. If NRI-values are predictors, the models are build only with one index as predictor instead of all available indices. In this case, only one predictor and one response variable is currently allowed. See help pages for 1m and g1m for any additional information. Note that this function does not store the entire information returned from a normal (g)lm-model. To get full (g)lm-models use either the function nri_best_performance to return best performing model(s) or extract nri-values with getNRI and build directly the model from respective index.

See details in Nri-plot-method for information about plotting.

Value

The function returns an object of class Nri. The list in the slot *multivariate* contains the new (g)lm information which depends on the kind of model which is applied:

1. lm.nri: The list contains the following items:

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• Estimate: Coefficient estimates for each index and term

• Std.Error: Standard errors

t.value: T-values
p.value: P-values
r.squared: R² values

2. glm.nri: The list contains the following items (depending on formula used):

• Estimate: Coefficient estimates for each index and term

• Std.Error: Standard errors

• t.value/z.value: T-values or Z-values

• p.value: P-values

Author(s)

Lukas Lehnert

See Also

```
plot, lm, glm, getNRI
```

Examples

hsdardocs

Load additional documents

Description

Access help documents and references for different methods.

Usage

hsdardocs(doc)

Arguments

doc

Name of document to load. Currently, only "References.pdf" and "Copyright" are available

48 hsdar_parallel

Author(s)

Lukas Lehnert

Examples

```
## Not run:
## Open references of hyperspectral vegetation indices (PDF-file)
hsdardocs("References.pdf")

## See copyrights of routines and data used in hsdar-package (ascii-file)
hsdardocs("Copyright")

## End(Not run)
```

hsdar_parallel

hsdar_parallel

Description

Get all functions which support parallel execution

Usage

```
hsdar_parallel()
```

Details

Parallel execution is performed via the **foreach**-package. Care is taken that a function will never run in parallel if the calling function is already using multicore processing.

Value

Vector containing supported function names

Author(s)

Lukas Lehnert

```
## Not run:
data(spectral_data)
## Load library
library(doMC)
## Register number of workers
registerDoMC(3)
supported_functions <- hsdar_parallel()
supported_functions</pre>
```

HyperSpecRaster 49

```
## Transform speclib using 3 cores
bd <- transformSpeclib(spectral_data)
## End(Not run)</pre>
```

HyperSpecRaster

Handle hyperspectral cubes using raster package (deprecated)

Description

The HyperSpecRaster-Class is deprecated. Use Speclib instead.

Usage

```
## S4 method for signature 'character, numeric'
HyperSpecRaster(x, wavelength, fwhm = NULL, SI = NULL, ...)
## S4 method for signature 'RasterLayer,numeric'
HyperSpecRaster(x, wavelength, fwhm = NULL, SI = NULL)
## S4 method for signature 'RasterBrick, numeric'
HyperSpecRaster(x, wavelength, fwhm = NULL, SI = NULL)
## S4 method for signature 'HyperSpecRaster'
HyperSpecRaster(x, wavelength)
## S4 method for signature 'Speclib, ANY'
brick(x, nrow, ncol, xmn, xmx, ymn, ymx, crs)
## S4 method for signature 'Speclib, ANY'
HyperSpecRaster(x, nrow, ncol, xmn, xmx, ymn, ymx, crs)
## S4 method for signature 'HyperSpecRaster, character'
writeStart(x, filename, ...)
## S4 method for signature 'HyperSpecRaster'
getValuesBlock(x, ...)
## S4 method for signature 'RasterLayer, Speclib'
writeValues(x, v, start)
## S4 method for signature 'RasterBrick, Speclib'
writeValues(x, v, start)
## S4 method for signature 'HyperSpecRaster, Speclib'
writeValues(x, v, start)
```

50 HyperSpecRaster

Arguments

X	Raster* object
wavelength	Vector containing wavelength for each band
fwhm	Optional vector containing full-width-half-max values. If length == 1 the same value is assumed for each band. Note that function does not check the integrity of the values
SI	Optional data.frame containing SI data
nrow	Optional. Number of rows in HyperspecRaster. If omitted, function will try to get the information from the SI in Speclib (attr(x, "rastermeta"))
ncol	Optional. Number of colums in HyperspecRaster. See nrow above.
xmn	Optional. Minimum coordinate in x-dimension. See nrow above.
xmx	Optional. Maximum coordinate in x-dimension. See nrow above.
ymn	Optional. Minimum coordinate in y-dimension. See nrow above.
ymx	Optional. Maximum coordinate in y-dimension. See nrow above.
crs	Optional. Object of class 'CRS' giving the coordinate system for HyperspecRaster. See nrow above.
	Additional arguments as for brick
filename	Name of file to create
V	Speclib or matrix of values
start	Integer. Row number (counting starts at 1) from where to start writing v

Value

HyperSpecRaster or RasterBrick

Author(s)

Lukas Lehnert

```
cellsize = c(1,1,1),
                                   cells.dim = c(cols, rows, 1)))
x <- SpatialPixelsDataFrame(grd, data = as.data.frame(spectra(spectra)))</pre>
## Write data to example file (example_in.tif) in workingdirectory
writeGDAL(x, fname = "example_in.tif", drivername = "GTiff")
## Examples for HyperSpecRaster using file example_in.tif
## Example 1:
## smoothing spectra
infile <- "example_in.tif"</pre>
outfile <- "example_result_1.tif"</pre>
wavelength <- spectra$wavelength</pre>
ra <- HyperSpecRaster(infile, wavelength)</pre>
tr <- blockSize(ra)</pre>
res <- writeStart(ra, outfile, overwrite = TRUE)</pre>
for (i in 1:tr$n)
  v <- getValuesBlock(ra, row=tr$row[i], nrows=tr$nrows[i])</pre>
  v <- smoothSpeclib(v, method="sgolay", n=25)</pre>
  res <- writeValues(res, v, tr$row[i])</pre>
}
res <- writeStop(res)</pre>
## Example 2:
## masking spectra and calculating vegetation indices
outfile <- "example_result_2.tif"</pre>
n_veg <- as.numeric(length(vegindex()))</pre>
res <- writeStart(ra, outfile, overwrite = TRUE, nl = n_veg)</pre>
for (i in 1:tr$n)
  v <- getValuesBlock(ra, row=tr$row[i], nrows=tr$nrows[i])</pre>
  mask(v) <- c(1350, 1450)
  v <- as.matrix(vegindex(v, index=vegindex()))</pre>
  res <- writeValues(res, v, tr$row[i])</pre>
res <- writeStop(res)</pre>
## End(Not run)
```

HyperSpecRaster* class HyperSpecRaster* class (deprecated)

Description

This is a deprecated class. Use Speclib-class instead.

idSpeclib

Details

Extension of *RasterBrick-class with three additional slots:

wavelength: A numeric vector giving the center wavelength for each band.

fwhm (optional): A numeric vector giving the full-width-half-max values for each band.

SI (optional): A data. frame containing additional information for each pixel.

The information in the three slots are used for the convertion to Speclib.

Author(s)

Lukas Lehnert

See Also

```
brick, Speclib
```

idSpeclib

Handling IDs of spectra

Description

Returning and setting ID of spectra in Speclib

Usage

```
idSpeclib(x)
idSpeclib(x) <- value</pre>
```

Arguments

x Object of class Speclib.

value Character vector of the same length as nspectra(x), or NULL.

Value

For idSpeclib<-, the updated object. Otherwise a vector giving the ID of each spectrum in Speclib is returned.

Author(s)

Lukas Lehnert

See Also

Speclib

import_USGS 53

Examples

```
data(spectral_data)
idSpeclib(spectral_data)
```

import_USGS

import USGS spectra

Description

Import and download spectral data from USGS spectral library

Usage

Arguments

url Character passing the url of the data. If NULL, the following URL is used:

'ftp://ftpext.cr.usgs.gov/pub/cr/co/denver/speclab/pub/spectral.library/splib06.library/ASCII/'

avl List of available files. Typically the result of USGS_get_available_files.

pattern Search pattern to define a subset of all available spectra.

retrieve Logical. Should the data be downloaded?

loadAsSpeclib Logical. If TRUE, an object of class "Speclib" is retured

tol Discrepancy of the wavelength values between different spectra.

Author(s)

Lukas Lehnert

```
## Not run:
## Retrieve all available spectra
avl <- USGS_get_available_files()

## Download all spectra matching "grass-fescue"
grass_spectra <- USGS_retrieve_files(avl = avl, pattern = "grass-fescue")
plot(grass_spectra)

## End(Not run)</pre>
```

54 makehull

makehull

Check continuum line

Description

Check if continuum line is intersecting the reflectance curve.

Usage

```
makehull(x, ispec)
```

Arguments

x Object of class Clman.

ispec Name or index of spectrum to be checked.

Value

Object of class list.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

```
transformSpeclib, addcp, deletecp, makehull, updatecl
Clman
```

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5),2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)

## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")

## Plot original line
par(mfrow = c(1,2))
plot(spec_clman, ispec = 1, subset = c(2480, 2500))

## Add fix point at 4595 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2495)

## Plot new line
plot(spec_clman, ispec = 1, subset = c(2480, 2500))

## Check new hull</pre>
```

mask 55

```
hull <- checkhull(spec_clman, 1)
hull$error

## Add fix point at 4596 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2496)

## Check new hull
hull <- checkhull(spec_clman, 1)
hull$error

hull <- makehull(spec_clman, 1)

## Transform spectra using band depth
spec_bd <- transformSpeclib(spec, method = "sh", out = "bd")

## Update continuum line of first spectrum
spec_bd <- updatecl(spec_bd, hull)

## Plot modified transformed spectrum
plot(spec_bd, FUN = 1)</pre>
```

mask

Mask spectra

Description

Returning and setting mask of spectra in Speclib. interpolate.mask linearly interpolates masked parts in spectra.

Usage

```
## $4 method for signature 'Speclib'
mask(object)
## $4 replacement method for signature 'Speclib,data.frame'
mask(object) <- value
## $4 replacement method for signature 'Speclib,list'
mask(object) <- value
## $4 replacement method for signature 'Speclib,numeric'
mask(object) <- value
## Linear interpolation of masked parts
interpolate.mask(object)</pre>
```

Arguments

object Object of class Speclib.

value Numeric vector, data frame or list giving the mask boundaries in wavelength

units. See details section.

56 mask

Details

Value may be an object of class vector, data frame or list. Data frames must contain 2 columns with the first column giving the lower (lb) and the second the upper boundary (ub) of the wavelength ranges to be masked. List must have two items consisting of vectors of length = 2. The first entry is used as lower and the second as upper boundary value. Vectors must contain corresponding lower and upper boundary values consecutively. The masked wavelength range(s) as defined by the lower and upper boundaries are excluded from the object of class Speclib.

Interpolation of masked parts is mainly intended for internal use. Interpolation is only possible if mask does not exceed spectral range of Speclib.

Value

For mask<-, the updated object. Otherwise a data frame giving the mask boundaries. interpolate.mask returns a new object of class Speclib.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

Speclib

```
data(spectral_data)
mask(spectral_data) ## NULL
## Mask from vector
spectral_data_ve <- spectral_data</pre>
mask(spectral_data_ve) <- c(1040,1060,1300,1450)
mask(spectral_data_ve)
## Mask from data frame
spectral_data_df <- spectral_data</pre>
mask(spectral_data_df) <- data.frame(lb=c(1040,1300),ub=c(1060,1450))</pre>
mask(spectral_data_df)
## Mask from list
spectral_data_li <- spectral_data</pre>
mask(spectral_data_li) <- list(lb=c(1040,1300),ub=c(1060,1450))</pre>
mask(spectral_data_li)
## Linear interpolation
plot(spectral_data)
plot(interpolate.mask(spectral_data_li), new=FALSE)
```

meanfilter 57

meanfilter

Apply mean filter

Description

Apply mean filter to data frame with spectra as rows and bands as columns. Filter size is passed as number of bands averaged at both sites of the respective band value.

Usage

```
meanfilter(spectra, p = 5)
```

Arguments

spectra Data frame containing spectra

p Filter size.

Value

Filtered data frame of same dimension as input data frame

Author(s)

Lukas Lehnert

See Also

```
smoothSpeclib
```

```
data(spectral_data)
spectra_filtered <- meanfilter(spectra(spectral_data), p = 10)
spectra(spectral_data) <- spectra_filtered</pre>
```

58 merge

merge

Merge speclibs

Description

Merge 2 Speclibs and their SI data

Usage

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

Arguments

x 1st Object of class Speclib to be merged.

y 2nd Object of class Speclib to be merged.

... Further arguments passed to generic functions. Currently ignored.

Value

Object of class Speclib.

Author(s)

Lukas Lehnert

See Also

```
Speclib
```

```
data(spectral_data)
sp1 <- spectral_data[c(1:10),]
sp2 <- spectral_data[c(11:20),]
speclib_merged <- merge(sp1, sp2)</pre>
```

nri 59

nri	Normalised ratio	index

Description

Calculate normalised ratio index (nri) for a single given band combination or for all possible band combinations. Calculating nri is a frequently used method to standardize reflectance values and to find relationships between properties of the objects and their spectral data.

Usage

```
nri(x, b1, b2, recursive = FALSE, bywavelength = TRUE)
```

Arguments

X	List of class Speclib or of class Nri for print and as.matrix methods.
b1	Band 1 given as band number or wavelength.
b2	Band 2 given as band number or wavelength.
recursive	If TRUE indices for all possible band combinations are calculated. If FALSE, only a single nri for the given bands in b1 and b2 is calculated.
bywavelength	Flag to determine if b1 and b2 are band number (bywavelength = FALSE) or wavelength (bywavelength = TRUE) values.
	Further arguments passed to generic functions. Currently ignored.

Details

Function performs the following calculation:

$$nri_{B1, B2} = \frac{R_{B1} - R_{B2}}{R_{B1} - R_{B2}};$$

with R being reflectance values at wavelength B1 and B2, respectively.

If recursive = TRUE, all possible band combinations are calculated.

Value

If recursive = FALSE, a data frame with index values is returned. Otherwise result is an object of class Nri. See glm.nri for applying a generalised linear model to an array of normalised ratio indices.

Author(s)

Lukas Lehnert

Nri-class

References

Sims, D.A.; Gamon, J.A. (2002). Relationships between leaf pigment content and spectral reflectance across a wide range of species, leaf structures and developmental stages. Remote Sensing of Environment: 81/2, 337 - 354.

Thenkabail, P.S.; Smith, R.B.; Pauw, E.D. (2000). Hyperspectral vegetation indices and their relationships with agricultural crop characteristics. Remote Sensing of Environment: 71/2, 158 - 182.

See Also

```
glm.nri, glm, Speclib, Nri
```

Examples

Nri-class

* Nri class

Description

Class to handle datasets containing normalized ratio indices of spectra.

Details

Object with slots:

- nri: Object of class DistMat3D containing nri values.
- fwhm: Vector or single numerical value giving the full-width-half-max value(s) for each band.
- wavelength: Vector with wavelength information.
- dimnames: Character vector containing band names used to calculate nri-values.
- multivariate: List defining the kind of test/model applied to the data and the model data. Only used after object has passed e.g. (g)lm.nri.
- SI: Data.frame containing additional data
- usagehistory: Vector giving information on history of usage of the object.

Nri-methods 61

Note

See figure in hsdar-package for an overview of classes in hsdar.

Author(s)

Lukas Lehnert

See Also

Speclib

Nri-methods

Methods for * Nri-class

Description

Methods to handle data in objects of class Nri.

Usage

```
## S4 method for signature 'Nri'
as.matrix(x, ..., named_matrix = TRUE)
## S4 method for signature 'Nri'
as.data.frame(x, na.rm = FALSE, ...)
## S4 method for signature 'Nri'
wavelength(object)
getFiniteNri(x)
```

Arguments

x, object List of class 'Nri'

na.rm Remove indices containing NA-values. Note that if TRUE, all indices are re-

moved which have at least one NA value.

named_matrix Flag if column and row names are set to band indices used for the calculation of

the nri-values.

... Further arguments passed to generic functions. Currently ignored.

Author(s)

Lukas Lehnert

See Also

```
glm.nri, glm, nri
```

62 nri_best_performance

```
nri_best_performance Best performing model(s) with NRI
```

Description

Get or mark best performing model(s) between narrow band indices and environmental variables

Usage

Arguments

nri	Object of class nri
glmnri	Object of class glmnri

n Number of models to return or mark

coefficient Name or index of coefficient to plot

predictor Name or index of term to plot

abs Use absolute value (e.g. for t-values)
findMax Find maximum or minimum values
best Output from nri_best_performance

uppertriang Flag to mark the upper triangle

Further arguments passed to glm function. These must be the same as used for

initial creation of glm.nri. For mark_nri_best_performance arguments are

passed to polygon.

Details

```
See details in glm.nri and glm.
```

Author(s)

Lukas Lehnert

See Also

```
glm.nri, glm
```

plot.Nri 63

Examples

plot.Nri

Plot function for (g)lm.nri and cor.test.nri

Description

Plot values in (generalised) linear modes and correlation tests from narrow band indices

Usage

```
## S4 method for signature 'Nri'
plot(x, coefficient = "p.value", predictor = 2,
    xlab = "Wavelength band 1 (nm)",
    ylab = "Wavelength band 2 (nm)", legend = TRUE,
    colspace = "hcl", col = c(10, 90, 60, 60, 10, 80),
    digits = 2, range = "auto", constraint = NULL,
    uppertriang = FALSE, ...)
```

Arguments

X	Object to be plotted.
coefficient	Name or index of coefficient to plot.
predictor	Name or index of term to plot.
xlab	Label for x-axis.
ylab	Label for y-axis.
legend	Flag if legend is plotted. If legend == "outer" the legend is plotted in the outer margins of the figure. This is useful if both diagonals are used.
colspace	Either "hcl" or "rgb". Colour space to be used for the plots.

plot.Nri

col	If colspace == "hcl", the vector is giving the minimum and maximum values of hue (element 1 & 2), chroma (element 3 & 4) and luminance (element 5 & 6). The optional element 7 is used as alpha value. See hcl for further explanation. If colspace == "rgb", a vector of length >= 2 giving the colours to be interpolated using colorRamp.
digits	Precision of labels in legend.
range	"auto" or a vector of length = 2 giving the range of values to be plotted.
constraint	A character string giving a constraint which values should be plotted. See examples section.
uppertriang	Flag if upper triangle is used for the plot. Note that if TRUE the current plot is used instead of starting a new plot
	Further arguments passed to plot.default.

Details

See details in glm. nri and glm.

Value

An invisible vector with minimum and maximum values plotted.

Author(s)

Lukas Lehnert

See Also

```
nri, glm.nri, glm, cor.test, t.test
```

plot.Specfeat 65

```
## Fit linear models between NRI-values and chlorophyll
lmnri <- lm.nri(nri_WV ~ chlorophyll, preddata = spec_WV)</pre>
## Plot r.squared
plot(lmnri)
## Example for EnMAP (Attention: Calculation time may be long!)
spec_EM <- spectralResampling(spectral_data, "EnMAP",</pre>
                               response_function = FALSE)
mask(spec_EM) <- c(300, 550, 800, 2500)
nri_EM <- nri(spec_EM, recursive = TRUE)</pre>
glmnri <- glm.nri(nri_EM ~ chlorophyll, preddata = spec_EM)</pre>
## Plot T values in lower and p-values in upper diagonal
## of the plot
## Enlarge margins for legends
par(mar = c(5.1, 4.1, 4.1, 5))
plot(glmnri, coefficient = "t.value", legend = "outer")
plot(glmnri, coefficient = "p.value", uppertriang = TRUE)
lines(c(400,1705),c(400,1705))
## End(Not run)
```

plot.Specfeat

Plot Specfeat

Description

Plot spectra in objects of class Specfeat. Specfeats contain spectral data after applying a transformation such as continuum removal (see function transformSpeclib.

Usage

Arguments

X	Object to be plotted
fnumber	Subscript of feature(s) to be plotted
stylebysubset	Name of column in SI table to be used for colour.
changecol	Flag indicating if line colours change according to values in coloumn defined by stylebysubset
changetype	Flag indicating if line types change according to values in coloumn defined by stylebysubset
autolegend	Flag if legend is plotted.
new	Flag if a new plot should be started.
	Further arguments passed to plot.default

plot.Speclib

Author(s)

Lukas Lehnert

See Also

```
nri, glm.nri, glm, cor.test, Nri-method, t.test, Nri-method, Specfeat
```

Examples

```
## Not run:
data(spectral_data)

## Transform speclib
bd <- transformSpeclib(spectral_data, method = "sh", out = "bd")

## Define features automatically
features <- define.features(bd)

##Example to isolate the features around 450nm, 700nm, 1200nm and 1500nm.
featureSelection <- specfeat(features, c(450,700,1200,1500))

## Plot features
plot(featureSelection, 1:4)

## Advanced plotting example
plot(featureSelection, 1:4, stylebysubset = "season")

plot(featureSelection, 1:4, stylebysubset = "season", changecol = FALSE, changetype = TRUE)

## End(Not run)</pre>
```

plot.Speclib

Plot speclib

Description

Plot Speclib in a new plot or adding it to an existing plot.

Usage

```
## S4 method for signature 'Speclib,ANY'
plot(x, FUN = NULL, new = TRUE, ...)
## S4 method for signature 'Clman,ANY'
plot(x, ispec, subset = NULL, numeratepoints = TRUE,
    hull.style = NULL, points.style = list(), ...)
```

plot.Speclib 67

Arguments

x Object of class Speclib	٥.
---------------------------	----

FUN Name of a function (character) or index or ID of single spectrum to plot (inte-

ger).

new If FALSE the plot is added to active existing plot.

ispec Subscript of spectrum to be plotted.

subset Vector of length = 2 containing minimum and maximum wavelength to plot.

numeratepoints Flag if continuum points are numerated and labeled.

hull.style List of arguments passed to lines to construct the continuum line.

points.style List of arguments passed to points to construct the continuum points. May be

NULL to suppress plotting of fix points.

... Further arguments passed to internal plot functions or to plot for objects of

class Speclib and Clman.

Details

The function may work in a couple of modes. The default way is to plot mean values (solid line) of all spectra and the standard deviations within bands. If data is assumed to be continuous the standard deviations are plotted as dashed lines otherwise error bars will indicate standard deviations.

The user has various options to change the way things are looking: With argument FUN the name of a function, the ID or the index of a certain spectrum may be specified. Note that if FUN is a function, this function will be applied to all spectra. If function should be applied to a subset of spectra, use function subset to define rules excluding certain spectra.

By passing a subset, the user may specify a spectral range to plot. Limits for x- and y-axis will be found automatically or may be passed separately.

Author(s)

Lukas Lehnert

See Also

Speclib

```
data(spectral_data)
## Set mask for channel crossing and water absorption bands
mask(spectral_data) <- c(1040, 1060, 1350, 1450)
## Simple example
plot(spectral_data, legend = list(x = "topleft"))
## Example with groups
plot(spectral_data, bygroups = TRUE, legend = list(x = "topleft"))</pre>
```

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```
## Example with function
par(mfrow = c(2,3))
plot(spectral_data, FUN = "min", main = "Minimum of speclib")
plot(spectral_data, FUN = "max", main = "Maximum of speclib")
plot(spectral_data, FUN = "median", main = "Median of speclib")
plot(spectral_data, FUN = "mean", main = "Mean of speclib")
plot(spectral_data, FUN = "var", main = "Variance of speclib")
```

predictHyperspec

Prediction based on train-object and Speclib

Description

Perform predictions based on a train-object from the **caret**-package and a hyperspectral dataset from **hsdar**. See help file to function predict.train of the **caret**-package for general information on prediction with **caret**.

Usage

```
## S4 method for signature 'train,.CaretHyperspectral,missing'
predictHyperspec(object, newdata, preProcess, ...)
## S4 method for signature 'train,.CaretHyperspectral,function'
predictHyperspec(object, newdata, preProcess, ...)
```

Arguments

object Object of class train from caret-package newdata Object of class Speclib or Nri to predict on.

preProcess Optional function to be applied on newdata prior to the prediction.

... Further arguments passed to original train function and/or to the preProcess-

function.

Value

Depending on the settings either a vector of predictions if type = "raw" or a data frame of class probabilities for type = "prob". In the latter case, there are columns for each class. For predict.list, a list results. Each element is produced by predict.train. See details in predict.train in the caret-package.

Author(s)

Lukas Lehnert

See Also

```
predict.train, Speclib
```

predictHyperspec 69

```
## Not run:
## The following example is taken from the journal paper
## "Hyperspectral Data Analysis in R: the hsdar Package"
## under review at the "Journal of Statistical Software"
data(spectral_data)
spectral_data <- smoothSpeclib(spectral_data, method = "sgolay", p = 15)</pre>
## Convert the chlorophyll measurements stored in the SI dataframe
## from SPAD-values into mg.
SI(spectral_data)$chlorophyll <-
  (117.1 * SI(spectral_data)$chlorophyll) /
  (148.84 - SI(spectral_data)$chlorophyll)
## Mask spectra
spectral_data <- spectral_data[, wavelength(spectral_data) >= 310 &
  wavelength(spectral_data) <= 1000]</pre>
## Transform reflectance values into band depth applying a segmented upper hull
## continuum removal.
spec_bd <- transformSpeclib(spectral_data, method = "sh", out = "bd")</pre>
## Search for fix points in the hull and define the parts of the spectra between
## two fix points as absorption "features"
features <- define.features(spec_bd)</pre>
## Select the chlorophyll absorption features at 460 nm and 670 nm for further
## processing
featureSpace <- specfeat(features, c(460, 670))</pre>
## Calculate all parameters from both selected features such as area, distance
## to Gauss curve etc.
featureSpace <- feature_properties(featureSpace)</pre>
## Set response and additional predictor variables for random forest model
featureSpace <- setResponse(featureSpace, "chlorophyll")</pre>
featureSpace <- setPredictor(featureSpace,</pre>
  names(SI(featureSpace))[4:ncol(SI(featureSpace))])
## Define training and cross validation for random forest model tuning
ctrl <- trainControl(method = "repeatedcv", number = 10, repeats = 5)</pre>
## Partition data set for training and validation
training_validation <- createDataPartition(featureSpace)</pre>
## Train random forest model based on training-subset
rfe_trained <- train(featureSpace[training_validation$Resample1,],</pre>
                      trainControl = ctrl, method = "rf")
## Predict on the validation data set
```

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PROSAIL

Simulate canopy spectrum

Description

Simulate a canopy spectrum using PROSAIL 5B

Usage

```
PROSAIL(N = 1.5, Cab = 40, Car = 8, Cbrown = 0.0, Cw = 0.01, Cm = 0.009, psoil = 0, LAI = 1, TypeLidf = 1, lidfa = -0.35, lidfb = -0.15, hspot = 0.01, tts = 30, tto = 10, psi = 0, parameterList = NULL, rsoil = NULL)
```

Arguments

N Structure parameter
 Cab Chlorophyll content
 Car Carotenoid content
 Cbrown Brown pigment content
 Cw Equivalent water thickness

Cm Dry matter content
psoil Dry/Wet soil factor
LAI Leaf area index

TypeLidf Type of leaf angle distribution. See details section

lidfa Leaf angle distribution. See details sectionlidfb Leaf angle distribution. See details section

hspot Hotspot parameter
tts Solar zenith angle
tto Observer zenith angle

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psi	Relative azimuth angle
parameterList	An optional object of class 'data.frame'. Function will iterate over rows of parameterList setting missing entries to default values. See examples section.
rsoil	An optional object of class 'Speclib' containing the background (soil) reflectance. Note that reflectance values must be in range [01].

Details

This function uses the FORTRAN code of PROSAIL model (Version 5B). For a general introduction see following web page and the links to articles provided there:

http://teledetection.ipgp.jussieu.fr/prosail/

The following table summarises the abbreviations of parameters and gives their units as used in PROSAIL. Please note that default values of all parameters were included with the intention to provide an easy access to the model and should be used with care in any scientific approach!

Parameter	Description of parameter	Units
N	Leaf structure parameter	NA
Cab	Chlorophyll a+b concentration	μ g/cm 2
Car	Carotenoid concentration	μ g/cm 2
Caw	Equivalent water thickness	cm
Cbrown	Brown pigment	NA
Cm	Dry matter content	g/cm ²
LAI	Leaf Area Index	NA
psoil	Dry/Wet soil factor	NA
hspot	Hotspot parameter	NA
tts	Solar zenith angle	deg
tto	Observer zenith angle	deg
psi	Relative azimuth angle	deg

Functions for distribution of leaf angles within the canopy may work in two modes, which is controlled via TypeLidf:

1. TypeLidf == 1 (default): lidfa is the average leaf slope and lidfb describes bimodality of leaf distribution. The following list gives an overview on typical settings:

LIDF type	lidfa	lidfb
Planophile	1	0
Erectophile	-1	0
Plagiophile	0	-1
Extremophile	0	1
Spherical (default)	-0.35	-0.15

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2. TypeLidf != 1: lidfa is the average leaf angle in degree (0 = planophile / 90 = erectophile); lidfb is 0

Value

An object of class Speclib. If parameterList is used, the parameter are stored in SI table of Speclib.

Note

The function is based on the FORTRAN version of the PROSAIL-code initially developed by Stephane JACQUEMOUD, Jean-Baptiste FERET, Christophe FRANCOIS and Eben BROADBENT. SAIL component has been developed by Wout VERHOEF.

Author(s)

Lukas Lehnert

References

Jacquemoud, S., Verhoef, W., Baret, F., Bacour, C., Zarco-Tejada, P.J., Asner, G.P., Francois, C., and Ustin, S.L. (2009): PROSPECT + SAIL models: a review of use for vegetation characterization, Remote Sensing of Environment, 113, S56-S66.

See Also

```
PROSPECT, Speclib
```

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PROSPECT	Simulate plant spectrum	

Description

Simulate plant spectrum using PROSPECT 5b or PROSPECT D. The inversion uses the concept after Feret et al. (2008) and uses PROSPECT 5B.

Usage

Arguments

N	Structure parameter	
Cab	Chlorophyll content	
Car	Carotenoid content	
Anth	Anthocyanin content	
Cbrown	Brown pigment content	
Cw	Equivalent water thickness	
Cm	Dry matter content	
transmittance	Logical flag, if transmittance instead of reflectance values are returned.	
parameterList	An optional object of class 'data.frame'. Function will iterate over rows of parameterList setting missing entries to default values. See examples section.	
version	Sets the version of PROSPECT to be used (either "5B" or "D").	
x, transmittance_spectra		
	Speclib(s) containing the reflectance/transmittance values to be simulated during inversion of PROSPECT.	
P0	Initial set of parameters (N, Cab etc.).	
sam	Logical if spectral angle mapper is used as distance measurement. If FALSE, the root mean square error is used. Note that this flag has only an effect if no transmittance spectra are passed.	

Parameters passed to anms from the **pracma**-package

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Details

This function uses the FORTRAN code of PROSPECT model (Version 5B an D). For a general introduction see following web page and the links to articles provided there:

```
http://teledetection.ipgp.jussieu.fr/prosail/
```

The following table summarises the abbreviations of parameters and gives their units as used in PROSPECT. Please note that default values of all parameters were included with the intention to provide an easy access to the model and should be used with care in any scientific approach!

Parameter	Description of parameter	Units
N	Leaf structure parameter	NA
Cab	Chlorophyll a+b concentration	μ g/cm 2
Car	Carotenoid concentration	μ g/cm 2
Anth	Anthocyanin content	μ g/cm 2
Cw	Equivalent water thickness	cm
Cbrown	Brown pigment	NA
Cm	Dry matter content	g/cm^2

The inversion uses the function arms from the **pracma**-package and implements the Matlab-Code developed by Feret et al. (2008). Please note that the inversion currently only uses version 5B.

Value

An object of class Speclib.

Note

The function is based on the FORTRAN version of the PROSPECT-code initially developed by Jean-Baptiste FERET, Stephane JACQUEMOUD and Christophe FRANCOIS.

Author(s)

Lukas Lehnert

References

Jacquemoud, S. and Baret, F. (1990). PROSPECT: A model of leaf optical properties spectra, Remote Sensing of Environment 34: 75 - 91.

Feret J.B., Francois C., Asner G.P., Gitelson A.A., Martin R.E., Bidel L.P.R., Ustin S.L., le Maire G., & Jacquemoud S. (2008), PROSPECT-4 and 5: advances in the leaf optical properties model separating photosynthetic pigments. Remote Sensing of Environment, 112, 3030-3043.

Feret J.B., Gitelson A.A., Noble S.D., & Jacquemoud S. (2017), PROSPECT-D: towards modeling leaf optical properties through a complete lifecycle, Remote Sensing of Environment, 193, 204-215.

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See Also

```
PROSAIL, anms, Speclib
```

Examples

```
## Single spectrum
spectrum <- PROSPECT(N = 1.3, Cab = 30, Car = 10, Cbrown = 0,</pre>
                     Cw = 0.01, Cm = 0.01)
plot(spectrum)
## Example using parameterList
## Test effect of leaf structure and chlorophyll content on
parameter \leftarrow data.frame(N = c(rep.int(seq(0.5, 1.5, 0.5), 2)),
                         Cab = c(rep.int(40, 3), rep.int(20, 3)))
spectra <- PROSPECT(parameterList = parameter)</pre>
## Print SI table
SI(spectra)
## Plot spectra for range from 400 to 800 nm
spectra <- spectra[,wavelength(spectra) >= 400 &
                    wavelength(spectra) <= 800]</pre>
plot(subset(spectra, Cab == 20), col = "red", ylim = c(0, 0.5))
plot(subset(spectra, Cab == 40), col = "green", new = FALSE)
```

Raster-methods

Rasterbased methods for spectra

Description

Methods to manipulate and save spectra in Speclibs stored as RasterBrick

Usage

```
## S4 method for signature 'Speclib,ANY'
extract(x, y, ...)
## S4 method for signature 'Speclib,character'
writeRaster(x, filename, ...)
```

Arguments

X	Speclib with RasterBrick-object for spectra
У	Object of any valid type to define area to extract
filename	Output filename
	Additionaly arguments passed to basic funtions in the raster-package

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Value

Speclib

Author(s)

Lukas Lehnert

rastermeta

Create list containing rastermeta-information

Description

Create valid objects for slot rastermeta in Speclib.

Usage

```
rastermeta(x, dim, ext, crs)
```

Arguments

X	Optional. Object of one of the following classes: "Raster", "RasterBrick", "RasterStack", "HyperSpecRaster".
dim	Optional. Vector with length == 2. The first and second elements give the number of rows and columns, respectively.
ext	Optional. Object of class extent.
crs	Optional. Object of class CRS.

Value

List with following elements (in exactly this order!):

- dim: Vector with length == 2. The first and second elements give the number of rows and columns, respectively.
- ext: Object of class extent.
- crs: Object of class CRS.

Author(s)

Lukas Lehnert

See Also

Speclib, HyperSpecRaster

rededge 77

rededge	Red edge parameter	

Description

Derive red edge parameters from hyperspectral data

Usage

```
rededge(x, smooth = TRUE, round = FALSE, ...)
```

Arguments

x List of class Speclib
 smooth Logical indicating if spectral data should be smoothed. See details section.
 round Logical indicating if resulting wavelength position should be rounded.

... Further arguments passed to derivative.speclib

Details

Shape and location of the red edge are commonly described by four parameters:

- R0: minimum reflectance in the red spectrum
- $\lambda 0$: wavelength of the minimum reflectance
- λp : inflection point
- Rs: shoulder wavelength

The red edge parameters are calculated as proposed in Bach (1995) from the spectral area between 550 and 900 nm. $\lambda 0$ is calculated as the last root before the maximum value of the 2nd derivation. The minimum reflectance is the reflectance at ($\lambda 0$). The inflection point is the root of the 2nd derivative function between the maximum value and the minimum value. The shoulder wavelength is the first root beyond the minimum value of the 2nd derivation.

Optional smoothing is performed with

```
smoothSpeclib(x, method = "spline", n = round(nbands(x)/10,0))
```

prior to all calculations. Note that reflectance values returned by the rededge-function are original values and not the smoothed reflectances. This would not be the case, if already smoothed reflectance values are passed to rededge-function.

Value

A data frame containing parameters for each spectrum.

Author(s)

Hanna Meyer

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References

Bach, H. (1995): Die Bestimmung hydrologischer und landwirtschaftlicher Oberflaechenparameter aus hyperspektralen Fernerkundungsdaten. Muenchner Geographische Abhandlungen Reihe B, Band B21.

See Also

```
vegindex, derivative.speclib, smoothSpeclib
```

Examples

```
data(spectral_data)
rd <- rededge(spectral_data)
boxplot(rd$R0 ~ SI(spectral_data)$season, ylab = "R0")</pre>
```

response_functions

Satellite sensor response functions

Description

Satellite sensor response functions for all sensor channels

Format

An object of class 'data.frame'

Details

Please do not access this data directly, since it contains only the response values without any spectral information.

Note

This data is kindly provided by operators of satellites. See hsdardocs("Copyright") for copyright information on spectral response functions.

- Quickbird: Copyright by DigitalGlobe, Inc. All Rights Reserved
- RapidEye: Copyright by RapidEye AG
- WorldView-2: Copyright by DigitalGlobe, Inc. All Rights Reserved

Handling supplementary information (SI) of spectra

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Description

Supplementary information (SI) can be any additional data available for each spectrum in a Speclibor Nri-object. These functions are used to set or return SI-data of a Speclib or Nri-object. Note that SI-data is automatically subsetted if indexing and extracting single spectra from a Speclib- or Nri-object. SI-data may encompass (several) raster files which must have the same extent, resolution and x- and y-dimensions as the raster file used as spectral information.

Usage

```
## S4 method for signature 'Speclib'
SI(object)

## S4 method for signature 'Speclib, ANY, ANY'
SI(object, i, j)

## S4 replacement method for signature 'Speclib, data.frame'
SI(object) <- value

## S4 replacement method for signature 'Speclib, matrix'
SI(object) <- value

## S4 method for signature 'Nri'
SI(object)

## S4 replacement method for signature 'Nri, data.frame'
SI(object) <- value

## S4 replacement method for signature 'Nri, matrix'
SI(object) <- value</pre>
```

Arguments

object	Object of class Speclib or Nri.
i	Index of rows to keep. Note that in combination with raster files in the SI, it is MUCH faster to pass row index instead of cutting the resulting data frame. Thus, SI(object, i) should be used instead of SI(object)[i,].
j	Index of columns to keep. See comment above for usage with raster files in the SI.
value	Data frame with nrow(value) == nspectra(object), NULL or vector with length nspectra(object). Alternatively, objects of class RasterLayer are accepted. Please note that the function does not check for integrity in the latter

SI

SI

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case (e.g., no error will occur if number of spectra does not match number of pixel in the RasterLayer-object).

Details

Names of items in SI are used within the function subset to select/deselect spectra via logical expression. Values can be accessed via the "\\$"-sign (see examples).

Value

For SI<-, the updated object. SI returns a data frame with SI data.

Author(s)

Lukas Lehnert

See Also

```
Speclib, Nri
```

Examples

```
data(spectral_data)
## Returning SI
si_spec <- SI(spectral_data)</pre>
head(si_spec)
## Adding new SI item
SI(spectral_data)$MeasurementID <- c(1:nspectra(spectral_data))</pre>
head(SI(spectral_data))
## Replacing SI
SI(spectral_data) <- SI(spectral_data)[,c(1:3)]</pre>
head(SI(spectral_data))
## Example for raster file in SI
## Create raster file using PROSAIL
## Run PROSAIL
parameter <- data.frame(N = c(rep.int(seq(0.5, 1.4, 0.1), 6)),
                         LAI = c(rep.int(0.5, 10), rep.int(1, 10),
                                  rep.int(1.5, 10), rep.int(2, 10),
                                  rep.int(2.5, 10), rep.int(3, 10)))
spectra <- PROSAIL(parameterList = parameter)</pre>
## Create SpatialPixelsDataFrame and fill data with spectra from
## PROSAIL
rows <- round(nspectra(spectra)/10, 0)</pre>
cols <- ceiling(nspectra(spectra)/rows)</pre>
grd <- SpatialGrid(GridTopology(cellcentre.offset = c(1,1,1),</pre>
                                  cellsize = c(1,1,1),
```

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```
cells.dim = c(cols, rows, 1)))
x <- SpatialPixelsDataFrame(grd,</pre>
                               data = as.data.frame(spectra(spectra)))
## Write data to example file (example_in.tif) in workingdirectory
writeGDAL(x, fname = "example_in.tif", drivername = "GTiff")
infile <- "example_in.tif"</pre>
wavelength <- wavelength(spectra)</pre>
ra <- speclib(infile, wavelength)</pre>
tr <- blockSize(ra)</pre>
## Write LAI to separate raster file
LAI <- SI(spectra)$LAI
SI_file <- "example_SI.tif"</pre>
SI_raster <- setValues(raster(infile), LAI)</pre>
SI_raster <- writeRaster(SI_raster, SI_file)</pre>
## Read LAI file and calculate NDVI for each pixel where LAI >= 1
outfile <- "example_result_ndvi.tif"</pre>
SI(ra) <- raster(SI_file)</pre>
names(SI(ra)) <- "LAI"</pre>
res <- writeStart(ra, outfile, overwrite = TRUE, nl = 1)</pre>
for (i in 1:tr$n)
  v <- getValuesBlock(ra, row=tr$row[i], nrows=tr$nrows[i])</pre>
  mask(v) <- c(1350, 1450)
  LAI <- SI(v)$LAI
  v <- as.matrix(vegindex(v, index="NDVI"))</pre>
  v[LAI \leftarrow 1] \leftarrow NA
  res <- writeValues(res, v, tr$row[i])</pre>
}
res <- writeStop(res)</pre>
```

smgm

SMGM

Description

Calculate Gaussian model on soil spectra

Usage

```
smgm(x, percentage = TRUE, gridsize = 50)
```

Arguments

x Object of class Speclib.

percentage Flag if spectra in x are in range [0, 100]. If FALSE, the spectra are scaled to

[0,100].

gridsize Size of the grid used to perform least squares approximation.

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Details

The algorithm fits a Gaussian function to the continuum points of the spectra in the spectral region between approx. 1500 to 2500 nm. The continuum points are derived constructing the convex hull of the spectra (see transformSpeclib). The Gaussian function requires three parameter: (1) the mean values which is set to the water fundamental of 2800 nm, (2) the absorption depth at 2800 nm, and (3) the distance to the inflection point of the function. The latter two parameters are iteratively chosen using a grid search. The mesh size of the grid can be adjusted with the gridsize parameter. Note that the function requires the spectral reflectance values to be in interval [0, 100].

Value

Object of class Speclib containing the fitted Gaussian spectra and the parameters derived from the Gaussian curve. The three parameters (absorption depth, R0; distance to the inflection point, sigma; area between the curve and 100 % reflectance, area) are stored in the SI of the new Speclib. Additionally, the function returns the final root mean square error of the Gaussian fit.

Note

The code is based on the IDL functions written by Michael L. Whiting.

Author(s)

Lukas Lehnert

References

Whiting, M. L., Li, L. and Ustin, S. L. (2004): Predicting water content using Gaussian model on soil spectra. Remote Sensing of Environment, 89, 535-552.

See Also

```
soilindex, Speclib
```

Examples

```
## Use PROSAIL to simulate spectra with different soil moisture content
Spektr.lib <- smoothSpeclib(PROSAIL(parameterList = data.frame(psoil = seq(0,1,0.1), LAI = 0)))
smgm_val <- smgm(Spektr.lib)
for (i in 1:nspectra(smgm_val))
   plot(smgm_val, FUN = i, new = i==1, col = i)
SI(smgm_val)</pre>
```

smoothSpeclib 83

smoothSpeclib	Smooth spectra	
---------------	----------------	--

Description

Smooth spectra using Savitzky-Golay filtering, lowess-, spline-functions or mean filter.

Usage

```
smoothSpeclib(x, method = "mean", ...)
```

Arguments

x Object of class Speclib.
 method Character string giving the method to be used. Predefined valid options are "sgolay", "lowess", "spline" and "mean". However, method can also be the (character)

ter) name of any other filter function (see examples).

Further arguments passed to filter functions. See examples.

Details

This function allows filtering using four different methods:

- Savitzky-Golay: Smoothing applying Savitzky-Golay-Filter. See sgolayfilt for details.
- Lowess: Smoothing applying lowess-Filter. See lowess for details.
- Spline: Smoothing applying spline-Filter. See spline for details.
- Mean: Smoothing applying mean-Filter. See meanfilter for details.

Value

Object of class Speclib.

Author(s)

Lukas Lehnert

References

Tsai, F. & Philpot, W. (1998): Derivative analysis of hyperspectral data. Remote Sensing of Environment 66/1. 41-51.

See Also

```
sgolayfilt, lowess, spline, meanfilter
```

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Examples

```
data(spectral_data)
## Example of predefined filter functions
## Savitzky-Golav
sgolay <- smoothSpeclib(spectral_data, method="sgolay", n=25)</pre>
spline <- smoothSpeclib(spectral_data, method="spline",</pre>
                          n=round(nbands(spectral_data)/10,0))
## Lowess
lowess <- smoothSpeclib(spectral_data, method="lowess", f=.01)</pre>
## Mean
meanflt <- smoothSpeclib(spectral_data, method="mean", p=5)</pre>
par(mfrow=c(2,2))
plot(spectral_data, FUN=1, main="Savitzky-Golay")
plot(sgolay, FUN=1, new=FALSE, col="red", lty="dotted")
plot(spectral_data, FUN=1, main="Spline")
plot(spline, FUN=1, new=FALSE, col="red", lty="dotted")
plot(spectral_data, FUN=1, main="Lowess")
plot(lowess, FUN=1, new=FALSE, col="red", lty="dotted")
plot(spectral_data, FUN=1, main="Mean")
plot(meanflt, FUN=1, new=FALSE, col="red", lty="dotted")
## Example of a not predefined filter function (Butterworth filter)
bf <- butter(3, 0.1)</pre>
bf_spec <- smoothSpeclib(spectral_data, method="filter", filt=bf)</pre>
plot(spectral_data, FUN=1, main="Butterworth filter")
plot(bf_spec, FUN=1, new=FALSE, col="red", lty="dotted")
```

soilindex

soilindex

Description

Function calculates a variety of hyperspectral soil indices

Usage

```
soilindex(x, index, returnHCR = "auto", weighted = TRUE, ...)
```

Arguments

x Object of class Speclib

index Character string. Name or definition of index or vector with names/definitions

of indices to calculate. See Details section for further information.

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returnHCR	If TRUE, the result will be of class HyperSpecRaster, otherwise it is a data frame. If "auto", the class is automatically determined by passed Speclib.
weighted	Logical indicating if reflectance values should be interpolated to fit wavelength position. If FALSE the reflectance values of nearest neighbour to passed position are returned. See get_reflectance for further explanation.
•••	Further arguments passed to derivative functions. Only used for indices requiring derivations.

Details

Index must be a charater vector containing pre-defined indices (selected by their name) or self defined indices or any combination of pre- and self-defined indices.

Pre-defined indices: The following indices are available:

Name	Formula	Reference*
BI_TM CI_TM	$((TM_1^2 + TM_2^2 + TM_3^2)/3)^{0.5**}$ $(TM_3 - TM_2)/(TM_3 + TM_2)**$	Mathieu et al. (1998) Escadafal and Huete (1991)
HI_TM NDI	$(2 \cdot TM_3 - TM_2 - TM_1)/(TM_2 - TM_1)**$ $(R_{840} - R_{1650})/(R_{840} + R_{1650})$	Escadafal et al. (1994) McNairn, H. and Protz, R. (1993)
NSMI RI	$(R_{1800} - R_{2119})/(R_{1800} + R_{2119})$ $R_{693}^2/(R_{447} \cdot R_{556}^3)$	Haubrock et al. (2008) Ben-Dor et al. (2006)
RI_TM	$TM_{693}/(TM_{447} \cdot R_{556})$ $TM_{3}^{2}/(TM_{1} \cdot TM_{2}^{3})**$	Madeira et al. (1997), Mathieu et al. (1998)
SI_TM SWIR SI	$(TM_3 - TM_1)/(TM_3 + TM_1)^{**}$ -41.59 · $(R_{2210} - R_{2090})$ + 1.24 · $(R_{2280} - R_{2090})$ + 0.64	Escadafal et al. (1994) Lobell et al. (2001)

^{*} For references please type: hsdardocs("References.pdf").

Self-defining indices:

Self-defined indices may be passed using the following syntax:

- Rxxx: Reflectance at wavelength 'xxx'. Note that R must be upper case.
- Dxxx: First derivation of reflectance values at wavelength 'xxx'. Note that D must be upper case.

Using this syntax, complex indices can be easily defined. Note that the entire definition of the index must be passed as one character string. Consequently, the NSMI would be written as (R1800-R2119)/(R1800+R2119)".

^{**} TM_1 denotes the first band of Landsat Thematic Mapper. Consequently, the hyperspectral data is resmapled to Landsat TM using spectralResampling prior to the calculation of the index. For resampling, the spectral response function is used.

specfeat specfeat

Value

A vector containing indices values. If index is a vector with length > 1, a data frame with ncol = length(index) and nrow = number of spectra in x is returned.

If function is called without any arguments, return value will be a vector containing all available indices in alphabetical order.

Author(s)

Lukas Lehnert

References

```
See hsdardocs("References.pdf")
```

See Also

```
vegindex, get_reflectance
```

Examples

```
data(spectral_data)
## Example calculating all available indices
## Get available indices
avl <- soilindex()
vi <- soilindex(spectral_data, avl)</pre>
```

specfeat

Function to isolate spectral features

Description

Function isolates specified absorption features previously identified by define.features.

Usage

specfeat 87

Arguments

x Object of class Speclib containing the band depth or ratio transformed re-

flectance spectra with additional information on feature limits calculated by

define. features. For plot this must be object of class specfeat.

FWL A vector containing one wavelength per feature to be isolated, e.g. the major

absorption features. Features which include these specified wavelengths will be

isolated.

fnumber Index of feature(s) to be plotted.

stylebysubset Name of variable to be used as grouping factor. May be selected from SI table,

groups or subgroups and must be convertible to factors.

changecol Flag, if line colour should be varied among groups changetype Flag, if line styles should be varied among groups

autolegend Flag if, legend is printed automatically.

new Create new plot or add data to existing one.

Further arguments passed to plot function.

Value

An object of class Specfeat containing the isolated features.

Author(s)

Hanna Meyer and Lukas Lehnert

See Also

```
define.features, cut_specfeat, Specfeat
```

Examples

```
data(spectral_data)
## Transform speclib
bd <- transformSpeclib(spectral_data, method = "sh", out = "bd")
## Define features automatically
features <- define.features(bd)

##Example to isolate the features around 450nm, 700nm, 1200nm and 1500nm.
featureSelection <- specfeat(features, c(450,700,1200,1500))

## Plot features
plot(featureSelection, fnumber = 1:4)

## Advanced plotting example
plot(featureSelection, fnumber = 1:4, stylebysubset = "season")</pre>
```

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Specfeat-class

* Specfeat class

Description

Class to handle spectral feature data.

Details

Class extends Speclib-class and adds two additional slots:

- features: List containing the spectra according to the features.
- featureLimits: List containing limits of features defined by define.features.

Note

See figure in hsdar-package for an overview of classes in hsdar.

Author(s)

Lukas Lehnert

See Also

Speclib

speclib

Methods to create objects of class Speclib

Description

Methods to create objects of class Speclib from various data types

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Usage

```
## S4 method for signature 'matrix, numeric'
speclib(spectra, wavelength, ...)
## S4 method for signature 'SpatialGridDataFrame, numeric'
speclib(spectra, wavelength, ...)
## S4 method for signature 'numeric, numeric'
speclib(spectra, wavelength, ...)
## S4 method for signature 'matrix,data.frame'
speclib(spectra, wavelength, ...)
## S4 method for signature 'SpatialGridDataFrame, data.frame'
speclib(spectra, wavelength, ...)
## S4 method for signature 'numeric,data.frame'
speclib(spectra, wavelength, ...)
## S4 method for signature 'matrix, matrix'
speclib(spectra, wavelength, ...)
## S4 method for signature 'SpatialGridDataFrame,matrix'
speclib(spectra, wavelength, ...)
## S4 method for signature 'numeric, matrix'
speclib(spectra, wavelength, ...)
## S4 method for signature 'hyperSpec, ANY'
speclib(spectra, wavelength, ...)
## S4 method for signature 'character, numeric'
speclib(spectra, wavelength, ...)
## S4 method for signature 'Speclib, numeric'
speclib(spectra, wavelength, ...)
## S4 method for signature 'Speclib'
print(x)
## S4 method for signature 'Speclib'
show(object)
createspeclib(spectra, wavelength, fwhm = NULL, SI = NULL,
        usagehistory = NULL, transformation = NULL,
        continuousdata = "auto", wlunit = "nm",
        xlabel = "Wavelength", ylabel = "Reflectance",
        rastermeta = NULL)
```

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is.speclib(x)

Arguments

spectra Data frame, matrix of raster object of class 'RasterBrick' or 'SpatialGridDataFrame'

with spectral data. Alternatively, spectra may be the path to a raster file contain-

ing hyperspectral data.

x, object Object to be converted to or from Speclib. For conversion to Speclib it can be

a of class 'data frame', 'matrix', 'list' or 'character string'. In the latter case x is interpreted as path to raster object and read by readGDAL. For

conversion from Speclib the object must be of class Speclib.

wavelength Vector with corresponding wavelength for each band. A matrix or data frame

may be passed giving the upper and lower limit of each band. In this case, the first column is used as lower band limit and the second as upper limit, respec-

tively.

fwhm Vector containing full-width-half-max values for each band

SI Data frame with supplementary information to each spectrum.

transformation Kind of transformation applied to spectral data (character). See transformSpeclib

for available ones. If transformation = NULL, no transformation is assumed

(default).

usagehistory Character string or vector used for history of usage

continuousdata Flag indicating if spectra are quasi continuous or discrete sensor spectra

wlunit Unit of wavelength in spectra

xlabel Label of wavelength data to be used for plots etc.
ylabel Label of spectral signal to be used for plots etc.

rastermeta List of meta information for SpatialGridDataFrame. If missing, meta data in

speclib is used. Use function rastermeta to create valid objects.

... Further arguments passed to specific (generic) functions or createspeclib

Details

See details in Speclib.

Value

An object of class Speclib containing the following slots is returned:

- wavelength: Vector with wavelength information
- fwhm: Vector or single numerical value giving the full-width-half-max value(s) for each band.
- spectra: Object of class '.Spectra' with three slots:
 - fromRaster: logical, indicating if spectral data is read from a RasterBrick-object.
 - spectra_ma: Matrix with ncol = number of bands and nrow = number. Used if fromRaster== FALSE

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spectra_ra: RasterBrick-object which is used if fromRaster == TRUE.

Contains reflectance, transmittance or absorbance values. Handle with function spectra.

- SI: Data frame containing additional data to each spectrum. May be used for linear regression etc. Handle with function SI.
- usagehistory: Vector giving information on history of usage of speclib. Handle with function usagehistory.
- rastermeta: List containing meta information to create *Raster objects from Speclib. Handle with function rastermeta.

Author(s)

Lukas Lehnert

See Also

```
Speclib, plot, readGDAL, mask,
idSpeclib, dim, spectra,
SI
```

Examples

```
data(spectral_data)
spectra <- spectra(spectral_data)
wavelength <- spectral_data$wavelength
spectra <- speclib(spectra,wavelength)</pre>
```

Speclib-class

* Speclib class

Description

Class to store and handle hyperspectral data in R

Details

Spectral data: The spectral data (usually reflectance values) are stored in an object of class '.Spectra'. This object may eiter contain the spectral data as a RasterBrick or as a matrix with columns indicating spectral bands and rows different samples, respectively. The Speclib-class provides converting routines to and from RasterBrick-class allowing to read and write geographic raster data via brick. Since R is in general not intended to be used for VERY large data sets, this functionality should be handled with care. If raster files are large, one should split them in multiple smaller ones and process each of the small files, separately. See the excellent tutorial 'Writing functions for large raster files' available on https://CRAN.R-project.org/package=raster and section '2.2.2 Speclibs from raster files' in 'hsdar-intro.pdf'.

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Spectral information: Specible contains wavelength information for each band in spectral data. This information is used for spectral resampling, vegetation indices and plotting etc. Since spectra can be handled either as continuous lines or as discrete values like for satellite bands, spectral information is handled in two principle ways:

- Continuous spectra: Data of spectrometers or hyperspectral (satellite) sensors. This data is plotted as lines with dotted lines indicating standard deviations by default.
- Non-continuous spectra: Data of multispectral satellite sensors. Here, data is plotted as solid
 lines and error bars at the mean position of each waveband indicating standard deviations by
 default.

The kind of data may be chosen by the user by setting the flag "continuousdata" (attr(x, "continuousdata")) or passing continuousdata = TRUE/FALSE, when initially converting data to Speclib-class. Take care of doing so, because some functions as spectralResampling may only work correctly with continuous data!

The unit of spectral data must be set initially, when converting data to speclib. Note that the package currently supports only "nm" as unit. This is particularly important for function like vegindex, which need to get correct bands out of the spectral data.

Technical description: An object of class Speclib contains the following slots:

- wavelength: Vector with wavelength information.
- fwhm: Vector or single numerical value giving the full-width-half-max value(s) for each band.
- spectra: Object of class '.Spectra' with three slots:
 - fromRaster: logical, indicating if spectral data is read from a RasterBrick-object.
 - spectra_ma: Matrix with ncol = number of bands and nrow = number. Used if fromRaster
 == FALSE
 - spectra_ra: RasterBrick-object which is used if fromRaster == TRUE.

Contains reflectance, transmittance or absorbance values. Handle with function spectra.

- SI: Data frame containing additional data to each spectrum. May be used for linear regression etc. Handle with function SI.
- usagehistory: Vector giving information on history of usage of speclib. Handle with function usagehistory.

Note

See figure in hsdar-package for an overview of classes in hsdar.

Author(s)

Lukas Lehnert

See Also

```
plot, readGDAL, mask, idSpeclib,
dim, spectra, SI
```

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```
speclib_raster-methods
```

Functions for processing of large hyperspectral raster files

Description

Functions for processing of large hyperspectral raster files using the low-level functions provided by the raster-package. For a detailed overview see the vignette "Writing functions for large raster files" shipped along with the raster-package.

Usage

```
## S4 method for signature 'Speclib'
blockSize(x)

## S4 method for signature 'Speclib, character'
writeStart(x, filename, ...)

## S4 method for signature 'Speclib'
getValuesBlock(x, ...)

## S4 method for signature 'Speclib, Speclib'
writeValues(x, v, start)

## S4 method for signature 'Speclib, matrix'
writeValues(x, v, start)

## S4 method for signature 'Speclib, numeric'
writeValues(x, v, start)
```

Arguments

Х	Object of class Speclib.
filename	Name of the new file to create.
V	Object to write the data to file. May be one of the following classes: "Speclib", "matrix" or "numeric".
start	Integer. Row number (counting starts at 1) from where to start writing v.
	Further arguements passed to respective functions in the raster-packages.

Author(s)

Lukas Lehnert

94 spectra

Description

Returning and setting spectra in Speclib

Usage

```
## S4 method for signature 'Speclib'
spectra(object, i, j, ...)

## S4 replacement method for signature 'Speclib,data.frame'
spectra(object) <- value

## S4 replacement method for signature 'Speclib,matrix'
spectra(object) <- value

## S4 replacement method for signature 'Speclib,numeric'
spectra(object) <- value

## S4 replacement method for signature 'Speclib,RasterBrick'
spectra(object) <- value</pre>
```

Arguments

object	Object of class Speclib.
i	Index of spectra to return. If missing all spectra are returned.
j	Index of bands to return. If missing all bands are returned.
•••	Passed to internal function. Currently only one parameter is accepted: return_names: Logical indicating, if names of columns and rows should be set to bandnames and idSpeclib.
value	Matrix or RasterBrick-object containing spectral values. If value is a matrix, columns are band values and rows are spectra.

Details

For spectra<-, the function does not check if dimensions of spectra match dimensions of Speclib. Additionally, no conversion into matrix is performed! If spectra are not correctly stored, errors in other functions may arise. Thus check always carefully, if spectra are modified by hand.

Value

For spectra<-, the updated object. Otherwise a matrix of the spectra in x is returned.

spectralResampling 95

Author(s)

Lukas Lehnert

See Also

Speclib

Examples

```
data(spectral_data)
## Manual plot of the first spectrum
plot(wavelength(spectral_data), spectra(spectral_data)[1,], type="l")
```

spectralResampling

Spectral resampling

Description

Resample spectra to (satellite) sensors

Usage

Arguments

x Object of class Speclib. Data to be spectrally resampled.

sensor Character or data.frame containing definition of sensor characteristics. See

details section for further information.

rm. NA If TRUE, channels which are not covered by input data wavelength are removed

continuousdata Definition if returned Speclib is containing continuous data or not.

response_function

If TRUE, the spectral response function of the sensor is used for integration, if

FALSE a Gaussian distribution is assumed and if NA the mean value of spectra[min(ch):max(ch)]

is calculated.

Details

The characteristics of (satellite) sensor to integrate spectra can be chosen from a list of already implemented sensors. See get.sensor.characteristics for available sensors.

Otherwise the characteristics can be passed as a data.frame with two columns: first column with lower bounds of channels and second column with upper bounds. Alternatively, the data.frame may encompass band centre wavelength and full-width-half-maximum values of the sensor. Function will check the kind of data passed by partially matching the names of the data frame: If any

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column is named "fwhm" or "center", it is assumed that data are band centre and full-width-half-maximum values.

If sensor characteristics are defined manually, a Gaussian response is always assumed.

Value

Object of class Speclib

Author(s)

Lukas Lehnert

See Also

```
get.sensor.characteristics, get.gaussian.response
```

Examples

```
## Load example data
data(spectral_data)
## Resample to RapidEye
data_RE <- spectralResampling(spectral_data, "RapidEye",</pre>
                               response_function = TRUE)
## Plot resampled spectra
plot(data_RE)
## Compare different methods of spectral resampling
par(mfrow=c(1,3))
ga <- spectralResampling(spectral_data, "RapidEye",</pre>
                          response_function = FALSE)
re <- spectralResampling(spectral_data, "RapidEye",</pre>
                          response_function = TRUE)
plot(re)
no <- spectralResampling(spectral_data, "RapidEye",</pre>
                          response_function = NA)
plot(no)
```

spectral_data

Hyperspectral samples

Description

Hyperspectral samples from a FACE experiment in Germany

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Usage

```
data(spectral_data)
```

Format

An object of class Speclib

Details

Data has been sampled during vegetation period 2014 in spring and summer. Measurements were taken with a HandySpec Field portable spectrometer (tec5 AG Oberursel, Germany). This device has two channels measuring incoming and reflected radiation simultaneously between 305 and 1705 nm in 1 nm steps.

Author(s)

Wolfgang A. Obermeier, Lukas Lehnert, Hanna Meyer

subset.nri

Subsetting Nri-objects

Description

Return subsets of Nri-objects which meet conditions.

Usage

```
## S4 method for signature 'Nri'
subset(x, subset, ...)
```

Arguments

x Object of class 'Nri'.

subset Logical expression indicating spectra to keep: missing values are taken as false.

See details section.

... Further arguments passed to agrep.

Details

Matchable objects are SI data. Use column names to identify the respectrive SI. See SI to access SI of a Nri. IDs of samples may be accessed using "id.nri" as variable name.

Value

Object of class Nri.

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Author(s)

Lukas Lehnert

See Also

Nri, SI

Examples

subset.speclib

Subsetting speclibs

Description

Return subsets of Speclibs which meet conditions.

Usage

```
## S4 method for signature 'Speclib'
subset(x, subset, ...)
```

Arguments

X	Object of class 'Speclib'.
subset	Logical expression indicating spectra to keep: missing values are taken as false. See details section.
	Further arguments passed to agrep.

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Details

Matchable objects are SI data. Use column names to identify the respective SI. See SI to access SI of a Speclib. IDs of spectra may be accessed using "id.speclib" as variable name. To subset certain wavelength ranges of a Speclib refer to mask.

Value

Object of class Speclib.

Author(s)

Lukas Lehnert

See Also

```
Speclib, SI, mask
```

Examples

```
data(spectral_data)
## Return names of SI data
names(SI(spectral_data))
## Devide into both seasons
sp_summer <- subset(spectral_data, season == "summer")
sp_spring <- subset(spectral_data, season == "spring")
## Plot both speclibs
plot(sp_summer, col="darkgreen")
plot(sp_spring, col="darkred", new=FALSE)</pre>
```

t.test

t-test for nri values

Description

Performs Student's t-tests for normalized ratio index values.

Usage

```
## S4 method for signature 'Nri'
t.test(x, ...)
```

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Arguments

```
x Object of class 'nri'.... Arguments to be passed to t.test.
```

Value

An object of class "data.frame"

Author(s)

Lukas Lehnert & Hanna Meyer

See Also

```
t.test, cor.test, Nri-method, Nri
```

Examples

transformSpeclib

Transform spectra

Description

Transform spectra by using convex hull or segmented upper hull

Usage

```
transformSpeclib(data, ..., method = "ch", out = "bd")
```

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Arguments

data Speclib to be transformed

method Method to be used. See details section.

out Kind of value to be returned. See details section.

... Further arguments passed to generic functions. Currently ignored.

Details

Function performs a continuum removal transformation by firstly establishing a continuum line/hull which connects the local maxima of the reflectance spectrum. Two kinds of this hull are well established in scientific community: the convex hull (e.g. Mutanga et al. 2004) and the segmented hull (e.g. Clark et al. 1987). Both hulls are established by connecting the local maxima, however, the precondition of the convex hull is that the resulting continuum line must be convex whereas considering the segmented hull it might be concave or convex but the algebraic sign of the slope is not allowed to change from the global maximum of the spectrum downwards to the sides. In contrast to a convex hull, the segmented hull is able to identify small absorption features.

Specify method = "ch" for the convex hull and method = "sh" for the segmented hull. The output might be "raw", "bd" or "ratio":

• "raw": the continuum line is returned

• "bd": the spectra are transformed to band depth by

$$BD_{\lambda} = 1 - \frac{R_{\lambda}}{CV_{\lambda}},$$

where BD is the band depth, R is the reflectance and CV is the continuum value at the wavelength λ .

• "ratio": the spectra are transformed by

$$BD_{\lambda} = \frac{R_{\lambda}}{CV_{\lambda}}.$$

In some cases it might be useful to apply smoothSpeclib before the transformation if too many small local maxima are present in the spectra. Anyway, a manual improvement of the continuum line is possible using addcp and deletecp.

Value

If out != "raw" an object of class Speclib containing transformed spectra is returned. Otherwise the return object will be of class Clman.

Author(s)

Hanna Meyer and Lukas Lehnert

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References

Clark, R. N., King, T. V. V. and Gorelick, N. S. (1987): Automatic continuum analysis of reflectance spectra. Proceedings of the Third Airborne Imaging Spectrometer Data Analysis Workshop, 30. 138-142.

Mutanga, O. and Skidmore, A. K. (2004): Hyperspectral band depth analysis for a better estimation of grass biomass (Cenchrus ciliaris) measured under controlled laboratory conditions International Journal of applied Earth Observation and Geoinformation, 5, 87-96.

See Also

```
Clman, addcp, deletecp, checkhull
```

Examples

```
data(spectral_data)

transformed_spectra <- transformSpeclib(spectral_data)

par(mfrow=c(1,2))
plot(spectral_data)
plot(transformed_spectra)</pre>
```

unmix

Unmix spectra

Description

Unmix spectra or spectra resampled to satellite bands using endmember spectra

Usage

```
unmix(spectra, endmember, returnHCR = "auto", scale = FALSE, ...)
```

Arguments

spectra	Input spectra of class 'speclib'
endmember	Endmember spectra of class 'speclib'
returnHCR	Set class of value. If TRUE, value will be of class 'HyperSpecRaster', otherwise a list is returned. If auto, function will switch to mode depending on input data characteristics.
scale	Flag to scale spectra to [0,1] if necessary.
• • •	Further arguments passed to $HyperSpecRaster$ (ignored if returnHCR = FALSE).

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Details

Linear spectral unmixing is a frequently used method to calculate fractions of land-cover classes (endmembers) within the footprint of pixels. This approach has originally been intended to be used for multispectral satellite images. The basic assumption is that the signal received at the sensor (ρ_{mix}) is a linear combination of n pure endmember signals (ρ_i) and their cover fractions (f_i) :

$$\rho_{mix} = \sum_{i=1}^{n} \rho_i f_i,$$

where $f_1, f_2, ..., f_n >= 0$ and $\sum_{i=1}^n f_i = 1$ to fulfill two constraints:

- 1. All fractions must be greater or equal 0
- 2. The sum of all fractions must be 1

Since this linear equation system is usually over-determined, a least square solution is performed. The error between the final approximation and the observed pixel vector is returned as vector (error) in list (returnSpatialGrid = FALSE) or as last band if returnSpatialGrid = TRUE.

Value

A list containing the fraction of each endmember in each spectrum and an error value giving the euclidean norm of the error vector after least square error minimisation.

Note

Unmixing code is based on "i.spec.unmix" for GRASS 5 written by Markus Neteler (1999).

Author(s)

Lukas Lehnert

References

Sohn, Y. S. & McCoy, R. M. (1997): Mapping desert shrub rangeland using spectral unmixing and modeling spectral mixtures with TM data. Photogrammetric Engineering and Remote Sensing, 63, 707-716

Examples

```
## Not run:
## Use PROSAIL to generate some vegetation spectra with different LAI
parameter <- data.frame(LAI = seq(0, 1, 0.01))
spectral_data <- PROSAIL(parameterList = parameter)

## Get endmember spectra
## Retrieve all available spectra
avl <- USGS_get_available_files()

## Download all spectra matching "grass-fescue"
grass_spectra <- USGS_retrieve_files(avl = avl, pattern = "grass-fescue")</pre>
```

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updatec1

Check continuum line

Description

Check if continuum line is intersecting the reflectance curve.

Usage

```
updatecl(x, hull)
```

Arguments

x Object of class Speclib transformed by transformSpeclib.hull Hull to be applied to x. Output of function makehull.

Value

Object of class Speclib.

Author(s)

Lukas Lehnert and Hanna Meyer

See Also

```
transformSpeclib, makehull, Speclib
```

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Examples

```
## Model spectra using PROSAIL
parameter <- data.frame(N = rep.int(c(1, 1.5), 2), LAI = c(1,1,3,3))
spec <- PROSAIL(parameterList=parameter)</pre>
## Transform spectra
spec_clman <- transformSpeclib(spec, method = "sh", out = "raw")</pre>
## Plot original line
par(mfrow = c(1,2))
plot(spec\_clman, ispec = 1, subset = c(2480, 2500))
## Add fix point at 4595 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2495)</pre>
## Plot new line
plot(spec\_clman, ispec = 1, subset = c(2480, 2500))
## Check new hull
hull <- checkhull(spec_clman, 1)</pre>
hull$error
## Add fix point at 4596 nm to continuum line of first spectrum
spec_clman <- addcp(spec_clman, 1, 2496)</pre>
## Check new hull
hull <- checkhull(spec_clman, 1)</pre>
hull$error
hull <- makehull(spec_clman, 1)</pre>
## Transform spectra using band depth
spec_bd <- transformSpeclib(spec, method = "sh", out = "bd")</pre>
## Update continuum line of first spectrum
spec_bd <- updatecl(spec_bd, hull)</pre>
## Plot modified transformed spectrum
plot(spec_bd, FUN = 1)
```

usagehistory

History of usage

Description

Function to read and write history of usage for Speclibs. Similar to a log file, the history of usage records processing steps applied to a Speclib.

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Usage

```
usagehistory(x)
usagehistory(x) <- value</pre>
```

Arguments

x Object of class Speclib

value Character string to be added to usagehistory or NULL, if usagehistory should be

deleted.

Value

For usagehistory<-, the updated object. Otherwise a vector containing the history of usage of Speclib is returned.

Author(s)

Lukas Lehnert

See Also

Speclib

Examples

```
data(spectral_data)
## Return history of usage
usagehistory(spectral_data)
## Deleting history of usage
usagehistory(spectral_data) <- NULL
spectral_data
## Adding entries
usagehistory(spectral_data) <- "New entry" ## Adding new entry
usagehistory(spectral_data) <- "New entry 2" ## Adding second entry
spectral_data</pre>
```

vegindex

vegindex

Description

Function calculates a variety of hyperspectral vegetation indices

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Usage

Arguments

X	Object of class Speclib
index	Character string. Name or definition of index or vector with names/definitions of indices to calculate. See Details section for further information.
returnHCR	If TRUE, the result will be of class HyperSpecRaster, otherwise it is a data frame. If "auto", the class is automatically determined by passed Speclib.
L	Factor for SAVI index. Unused for other indices.
weighted	Logical indicating if reflectance values should be interpolated to fit wavelength position. If FALSE the reflectance values of nearest neighbour to passed position are returned. See get_reflectance for further explanation.
• • •	Further arguments passed to derivative functions. Only used for indices requiring derivations.

Details

Index must be a charater vector containing pre-defined indices (selected by their name) or self defined indices or any combination of pre- and self-defined indices.

Pre-defined indices: The following indices are available:

Name	Formula	Reference*
Boochs	D_{703}	Boochs et al. (1990)
Boochs2	D_{720}	Boochs et al. (1990)
CAI	$0.5 \cdot (R_{2000} + R_{2200}) - R_{2100}$	Nagler et al. (2003)
CARI	$a = (R_{700} - R_{550})/150$	Kim et al. (1994)
	$b = R_{550} - (a \cdot 550)$	
	$R_{700} \cdot abs(a \cdot 670 + R_{670} + b)/R_{670}$	
	$(a^2+1)^{0.5}$	
Carter	R_{695}/R_{420}	Carter (1994)
Carter2	R_{695}/R_{760}	Carter (1994)
Carter3	R_{605}/R_{760}	Carter (1994)
Carter4	R_{710}/R_{760}	Carter (1994)
Carter5	R_{695}/R_{670}	Carter (1994)
Carter6	R_{550}	Carter (1994)
CI	$R_{675} \cdot R_{690} / R_{683}^2$	Zarco-Tejada et al.
		(2003)
CI2	$R_{760}/R_{700}-1$	Gitelson et al. (2003)
ClAInt	$\int_{600nm}^{735nm} R$	Oppelt and Mauser
	o dodinin	(2004)
CRI1	$1/R_{515} - 1/R_{550}$	Gitelson et al. (2003)

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CRI2	$1/R_{515} - 1/R_{770}$	Gitelson et al. (2003)
CRI3	$1/R_{515} - 1/R_{550} \cdot R_{770}$	Gitelson et al. (2003)
CRI4	$1/R_{515} - 1/R_{700} \cdot R_{770}$	Gitelson et al. (2003)
D1	D_{730}/D_{706}	Zarco-Tejada et al.
	,	(2003)
D2	D_{705}/D_{722}	Zarco-Tejada et al.
	1007 122	(2003)
Datt	$(R_{850} - R_{710})/(R_{850} - R_{680})$	Datt (1999b)
Datt2	R_{850}/R_{710}	Datt (1999b)
Datt3	D_{754}/D_{704}	Datt (1999b)
Datt4	$R_{672}/(R_{550} \cdot R_{708})$	Datt (1998)
Datt5		Datt (1998)
Datt6	$R_{672}/R_{550} = (R_{860})/(R_{550} \cdot R_{708})$	Datt (1998)
Datt7	$(R_{860} - R_{2218})/(R_{860} - R_{1928})$	Datt (1999a)
Datt8	$(R_{860} - R_{1788})/(R_{860} - R_{1928})$	Datt (1999a)
DD	$(R_{749} - R_{720}) - (R_{701} - R_{672})$	le Maire et al. (2004)
DDn	$2 \cdot (R_{710} - R_{660} - R_{760})$	le Maire et al. (2008)
DPI	$(D_{688} \cdot D_{710})/D_{697}^2$	Zarco-Tejada et al.
		(2003)
DWSI1	R_{800}/R_{1660}	Apan et al. (2004)
DWSI2	R_{1660}/R_{550}	Apan et al. (2004)
DWSI3	R_{1660}/R_{680}	Apan et al. (2004)
DWSI4	R_{550}/R_{680}	Apan et al. (2004)
DWSI5	$(R_{800} + R_{550})/(R_{1660} + R_{680})$	Apan et al. (2004)
EGFN	$(\max(D_{650:750}) - \max(D_{500:550}))/$	Penuelas et al.
	$(\max(D_{650:750}) + \max(D_{500:550}))$	(1994)
EGFR	$\max(D_{650:750})/\max(D_{500:550})$	Penuelas et al. (1994)
EVI	$2.5 \cdot ((R_{800} - R_{670}))$	Huete et al. (1997)
	$(R_{800} - (6 \cdot R_{670}) - (7.5 \cdot R_{475}) + 1))$	` ,
GDVI	$(R_{800}^n - R_{680}^n)/(R_{800}^n + R_{680}^n)^{**}$	Wu (2014)
GI	R_{554}/R_{677}	Smith et al. (1995)
Gitelson	$1/R_{700}$	Gitelson et al. (1999)
Gitelson2	$(R_{750} - R_{800}/R_{695} - R_{740}) - 1$	Gitelson et al. (2003)
GMI1	R_{750}/R_{550}	Gitelson et al. (2003)
GMI2	R_{750}/R_{700}	Gitelson et al. (2003)
Green NDVI	$(R_{800} - R_{550})/(R_{800} + R_{550})$	Gitelson et al. (1996)
LWVI_1		Galvao et al. (2005)
LWVI_1 LWVI_2	$(R_{1094} - R_{983})/(R_{1094} + R_{983})$ $(R_{1094} - R_{1205})/(R_{1094} + R_{1205})$	Galvao et al. (2005) Galvao et al. (2005)
		· · ·
Maccioni	$(R_{780} - R_{710})/(R_{780} - R_{680})$	Maccioni et al. (2001)
MCARI	$((R_{700} - R_{670}) - 0.2 \cdot (R_{700} - R_{550}))$	Daughtry et al. (2000)
MOADHOGAIH	(R_{700}/R_{670})	D 1 (2000)
MCARI/OSAVI	//B	Daughtry et al. (2000)
MCARI2	$((R_{750} - R_{705}) - 0.2 \cdot (R_{750} - R_{550}))$	Wu et al. (2008)
	(R_{750}/R_{705})	
MCARI2/OSAVI2	/ · ·	Wu et al. (2008)
mND705	$(R_{750} - R_{705})/(R_{750} + R_{705} - 2 \cdot R_{445})$	Sims and Gamon (2002)
mNDVI	$(R_{800} - R_{680})/(R_{800} + R_{680} - 2 \cdot R_{445})$	Sims and Gamon (2002)
MPRI	$(R_{515} - R_{530})/(R_{515} + R_{530})$	Hernandez-Clemente et al.

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		(2011)
mDEID	Rad adge inflaction point using Gaussein fit	(2011) Miller et al. (1000)
mREIP	Red-edge inflection point using Gaussain fit	Miller et al. (1990)
MSAVI	$0.5 \cdot (2 \cdot R_{800} + 1 - (2 \cdot R_{800} + 1)^2 + (2 \cdot R_{800} + 1)$	Qi et al. (1994)
MCI	$((2 \cdot R_{800} + 1)^2 - 8 \cdot (R_{800} - R_{670}))^{0.5})$	Ht and Dl- (1000)
MSI	R_{1600}/R_{817}	Hunt and Rock (1989)
mSR	$(R_{800} - R_{445})/(R_{680} - R_{445})$	Sims and Gamon (2002)
mSR2	$(R_{750}/R_{705}) - 1/(R_{750}/R_{705} + 1)^{0.5}$	Chen (1996)
mSR705	$(R_{750} - R_{445})/(R_{705} - R_{445})$	Sims and Gamon (2002)
MTCI	$(R_{754} - R_{709})/(R_{709} - R_{681})$	Dash and Curran (2004)
MTVI	$1.2 \cdot (1.2 \cdot (R_{800} - R_{550}) -$	Haboudane et al.
MDLI	$2.5 \cdot (R_{670} - R_{550}))$	(2004)
NDLI	$(log(1/R_{1754}) - log(1/R_{1680}))/$	Serrano et al. (2002)
MDM	$(log(1/R_{1754}) + log(1/R_{1680}))$	G 1 (2002)
NDNI	$(log(1/R_{1510}) - log(1/R_{1680}))/$	Serrano et al. (2002)
NIDI II	$(log(1/R_{1510}) + log(1/R_{1680}))$	T. 1 (1050)
NDVI	$(R_{800} - R_{680})/(R_{800} + R_{680})$	Tucker (1979)
NDVI2	$(R_{750} - R_{705})/(R_{750} + R_{705})$	Gitelson and Merzlyak
	(D. D.) ((D. D.)	(1994)
NDVI3	$(R_{682} - R_{553})/(R_{682} + R_{553})$	Gandia et al. (2004)
NDWI	$(R_{860} - R_{1240})/(R_{860} + R_{1240})$	Gao (1996)
NPCI	$(R_{680} - R_{430})/(R_{680} + R_{430})$	Penuelas et al. (1994)
OSAVI	$(1+0.16) \cdot (R_{800} - R_{670})/$	Rondeaux et al.
	$(R_{800} + R_{670} + 0.16)$	(1996)
OSAVI2	$(1+0.16) \cdot (R_{750} - R_{705})/$	Wu et al. (2008)
	$(R_{750} + R_{705} + 0.16)$	
PARS	R_{746}/R_{513}	Chappelle et al. (1992)
PRI	$(R_{531} - R_{570})/(R_{531} + R_{570})$	Gamon et al. (1992)
PRI_norm	$PRI \cdot (-1)/(RDVI \cdot R_{700}/R_{670})$	Zarco-Tejada et al.
		(2013)
PRI*CI2	PRI · CI2	Garrity et al. (2011)
PSRI	$(R_{678} - R_{500}/R_{750}$	Merzlyak et al. (1999)
PSSR	R_{800}/R_{635}	Blackburn (1998)
PSND	$(R_{800} - R_{470})/(R_{800} - R_{470})$	Blackburn (1998)
PWI	R_{900}/R_{970}	Penuelas et al. (1997)
RDVI	$(R_{800} - R_{670})/\sqrt{R_{800} + R_{670}}$	Roujean and Breon (1995)
REP_LE	Red-edge position through linear extrapolation.	Cho and Skidmore (2006)
REP_Li	$R_{re} = (R_{670} + R_{780})/2$	Guyot and Baret (1988)
	$700 + 40 \cdot ((R_{re} - R_{700})/(R_{740} - R_{700}))$	
SAVI	$(1+L)\cdot (R_{800}-R_{670})/(R_{800}+R_{670}+L)$	Huete (1988)
SIPI	$(R_{800} - R_{445})/(R_{800} - R_{680})$	Penuelas et al. (1995),
		Penuelas et al. (1995a)
SPVI	$0.4 \cdot 3.7 \cdot (R_{800} - R_{670}) - 1.2 \cdot$	Vincini et al. (2006)
	$((R_{530} - R_{670})^2)^{0.5}$	
SR	R_{800}/R_{680}	Jordan (1969)
SR1	R_{750}/R_{700}	Gitelson and Merzlyak
		(1997)
SR2	R_{752}/R_{690}	Gitelson and Merzlyak
		(1997)

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SR3	R_{750}/R_{550}	Gitelson and Merzlyak (1997)
SR4	R_{700}/R_{670}	McMurtey et al. (1994)
SR5	R_{675}/R_{700}	Chappelle et al. (1992)
SR6	R_{750}/R_{710}	Zarco-Tejada and Miller (1999)
SR7	R_{440}/R_{690}	Lichtenthaler et al. (1996)
SR8	R_{515}/R_{550}	Hernandez-Clemente et al. (2012)
SRPI	R_{430}/R_{680}	Penuelas et al. (1995)
SRWI	R_{850}/R_{1240}	Zarco-Tejada et al. (2003)
Sum_Dr1	$\sum_{\substack{i=626\\ \Sigma_{i=680}}}^{795} D1_i$	Elvidge and Chen (1995)
Sum_Dr2	$\sum_{i=0}^{i=020} D1_i$	Filella and Penuelas
_	<i>∠</i> 1=080 °	(1994)
SWIR FI	$R_{2133}^2/(R_{2225} \cdot R_{2209}^3)$	Levin et al. (2007)
SWIR LI	$3.87 \cdot (R_{2210} - R_{2090}) -$	Lobell et al. (2001)
	$27.51 \cdot (R_{2280} - R_{2090}) - 0.2$	
SWIR SI	$-41.59 \cdot (R_{2210} - R_{2090}) +$	Lobell et al. (2001)
	$1.24 \cdot (R_{2280} - R_{2090}) + 0.64$	
SWIR VI	$37.72 \cdot (R_{2210} - R_{2090}) +$	Lobell et al. (2001)
	$26.27 \cdot (R_{2280} - R_{2090}) + 0.57$	
TCARI	$3 \cdot ((R_{700} - R_{670}) - 0.2 \cdot (R_{700} - R_{550})$	Haboudane et al. (2002)
	$(R_{700}/R_{670}))$	
TCARI/OSAVI	TCARI/OSAVI	Haboudane et al. (2002)
TCARI2	$3 \cdot ((R_{750} - R_{705}) - 0.2 \cdot (R_{750} - R_{550}) \cdot$	Wu et al. (2008)
	$(R_{750}/R_{705}))$	
TCARI2/OSAVI2	TCARI2/OSAVI2	Wu et al. (2008)
TGI	$-0.5(190(R_{670} - R_{550}) - 120(R_{670} - R_{480}))$	Hunt et al. (2013)
TVI	$0.5 \cdot (120 \cdot (R_{750} - R_{550}) -$	Broge and Leblanc
	$200 \cdot (R_{670} - R_{550}))$	(2000)
Vogelmann	R_{740}/R_{720}	Vogelmann et al. (1993)
Vogelmann2	$(R_{734} - R_{747})/(R_{715} + R_{726})$	Vogelmann et al. (1993)
Vogelmann3	D_{715}/D_{705}	Vogelmann et al. (1993)
Vogelmann4	$(R_{734} - R_{747})/(R_{715} + R_{720})$	Vogelmann et al. (1993)

^{*} For references please type: hsdardocs("References.pdf").

Self-defining indices:

Self-defined indices may be passed using the following syntax:

- Rxxx: Reflectance at wavelength 'xxx'. Note that R must be upper case.
- Dxxx: First derivation of reflectance values at wavelength 'xxx'. Note that D must be upper case.

^{**} For GDVI n must be defined appending an underscore and the intended exponent to the index name. E.g., for n = 2, the correct index name would be "GDVI_2". Note that GDVI-indices with n = 2, 3, 4 will be derived if all available indices are calculated.

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Using this syntax, complex indices can be easily defined. Note that the entire definition of the index must be passed as one character string. Consequently, the NDVI would be written as "(R800-R680)/(R800+R680)".

Value

A vector containing indices values. If index is a vector with length > 1, a data frame with ncol = length(index) and nrow = number of spectra in x is returned.

If function is called without any arguments, return value will be a vector containing all available indices in alphabetical order.

Author(s)

Hanna Meyer and Lukas Lehnert

References

```
See hsdardocs("References.pdf")
```

See Also

```
soilindex, derivative.speclib, rededge, get_reflectance
```

Examples

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wavelength

Handling wavelength and fwhm

Description

Methods to access and set wavelength (band center) and full-width-half-max (fwhm) values for class Speclib.

Usage

```
## S4 method for signature 'Speclib'
wavelength(object)

## S4 replacement method for signature 'Speclib,data.frame'
wavelength(object) <- value

## S4 replacement method for signature 'Speclib,numeric'
wavelength(object) <- value

## S4 method for signature 'Speclib'
fwhm(object)

## S4 replacement method for signature 'Speclib,numeric'
fwhm(object) <- value</pre>
```

Arguments

object Object of class Speclib.

value Numeric vector or data.frame containing wavelength values.

Details

Wavelength (band center) and full-width-half-max (fwhm) values are given for each spectral band. The wavelength is mandatory for creation of Speclib and is used within the whole functionality of the package (e.g., smoothSpeclib, spectralResampling, vegindex, nri, plot, mask).

Value

For wavelength<- and fwhm<-, the updated object. Otherwise a numeric vector of the wavelength and fwhm-values is returned.

Author(s)

Lukas Lehnert

See Also

Speclib

wavelength 113

Examples

```
data(spectral_data)
wavelength(spectral_data)
```

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