

Package ‘uFTIR’

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Type Package

Title Process and Analyze Agilent Cary 620 FTIR Microscope Images

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Description A set of tools to read, process, and summarize Agilent Cary 620 uFTIR Microscope hyperspectral images primarily intended for microplastic analysis.

License GPL-3

Encoding UTF-8

LazyData true

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'mask_sam.r' 'matrix_sam.r' 'mosaic_chunk.R' 'mosaic_compose.R'
'mosaic_info.R' 'mosaic_sam.R' 'plot_methods.R' 'preprocess.R'
'profile_methods.R' 'rgdal_polygonize.R' 'sam_load.R'
'sam_write.r' 'smooth_sam.R' 'summary_methods.R'
'tile_base_corr.R' 'tile_read.R' 'tile_sam.R' 'uFTIR.R'
'wavealign.R'

SystemRequirements C++11, GDAL >= 1.11.4, PROJ.4 (proj >= 4.8.0)

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as.clipper

As Clipper

Description

Coerce a [SAM](#) or [Smooth](#) object to clipper. A function useful to call [highlight_substance](#) when clipping is not necessary.

Usage

```
as.clipper(object, slice = 1)
```

Arguments

`object` The [SAM](#) or [Smooth](#) object to be coerced.
`slice` The slice to keep (a clipper is essentially a matrix).

Value

and S3 clipper

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
x <- tile_sam(x)
x <- as.clipper(x)
```

<code>clipmask-class</code>	<i>Class holding info for the clipper</i>
-----------------------------	---

Description

The filters we used to support our samples under the uFTIR microscope are round. This means that a cube might not be the best representation of our data, as they have noisy borders that might be misinterpreted by the program. Then, I implemented a [clipper](#) function to clip objects of class [SAM](#) or [Smooth](#) (even an S3 matrix) before summarizing them. Although this can be done step by step, you can also create a clip-mask for the [summary_sam](#) function to clip them for you. To create this clip-mask, you can call the function [toClip](#) which returns an object of this class.

The class has yet another use. Since the samples are placed under the microscope by hand, the cropping area is not always the same and (usually) it has to be adjusted. To have a visual aid, you can use the function [toClip](#), to then plot a [SAM](#) or [Smooth](#) and overlay the cropping circle by calling [polygon](#) and the `xycoords` slot of an object of this class (returned by [toClip](#)). Using that process you can test manually different center points and radius for the clipping circle.

Value

An S4 object of class "clipmask" to be used (internally) by the [clipper](#) function.

Slots

`xycoords` `polygon` (circle) coordinates to plot. matrix with two columns (x,y).

`rad` circle radius. integer.

`centre` circle centre. Numeric vector of two elements (x,y).

See Also

[toClip](#) [clipper](#)

Examples

NULL

clipper

Clipper

Description

A function to clip a [SAM](#) or [Smooth](#) object object (also a matrix, but I recommed this only for debugging). ALthough you can use this function in a 'step by step' process it is otherwise directly called by [summary_sam](#) which uses a [clipmask](#) object (returned by [toClip](#) as a clipping mask).

Usage

```
clipper(tarjet, centre = c(128, 128), rad = 120, slice = 1)
```

Arguments

tarjet	object of class SAM or Smooth (or matrix) to clip.
centre	The coordinates of the polygon centre (x,y).
rad	The circle radius.
slice	which slice of the SAM or Smooth object should be clipped? (only one!)

Value

The tarjet object clipped as matrix (and S3 clipper -not for human consumption).

See Also

[toClip](#) [codeclipmask](#) [summary_sam](#).

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
x <- tile_sam(x)
x <- smooth_sam(x, as.integer(length(primpke@clusternames)), window = 3, 1)

clip <- toClip(8,20,c(10,10))
plot(x)
polygon(clip@xycoords)

x <- clipper(x, clip@centre, clip@rad, 1)
```

`get_profile`*Get profile*

Description

To check whether the program is doing a good job or not (an the library is appropriate), it is usually a good idea to recover the expectra of all the pixels that match a given cluster or substance to plot the spectra and compare it to a standard (the library entrance for that cluster/substance for example).

This function allows you to retrieve the spectra of all pixels that match a given cluster/substance and returns a matrix that can then be plotted to do the checking. It also allows the user to plot the object and overlay the points that match a feature. The latter can be achieved also by using the [highlight_substance](#) function. However, that function is only implemented for S3 "clipper" objects, so this might be a way to overcome that limitation.

`get_profile_all` will replace `get_profile_sinfo` some day. For now it does the same thing BUT it works only for clipper objects and you need to pass the fpa size and the wavenumber (waves) vector. It was designed to run in the cluster. It goes well with [sam_write](#), [sam_load](#).

Usage

```
get_profile(x, where, ...)  
  
## S4 method for signature 'togetprofile,Tile'  
get_profile(x, where, ...)  
  
## S4 method for signature 'togetprofile,list'  
get_profile(x, where, ...)  
  
## S4 method for signature 'clipper,character'  
get_profile(x, where, dst_cluster, fpa, waves)  
  
get_profile_tile(  
  x,  
  where,  
  dst_cluster,  
  plotpol = TRUE,  
  plotpt = FALSE,  
  cluster = TRUE,  
  slice = 1,  
  clusternames = NULL,  
  ...  
)  
  
get_profile_sinfo(  
  x,  
  where,  
  dst_cluster,
```

```

    plotpol = TRUE,
    plotpt = FALSE,
    cluster = TRUE,
    slice = 1,
    clusternames = NULL,
    ...
)

get_profile_all(x, where, dst_cluster, fpa, waves)

```

Arguments

x	Object of class sam , Smooth or clipper that hold the processed tile/mosaic indicated in where.
where	Original object processed. It might be of class Tile or SpectralInfo . However, in the second case, where should be a labeled list of two elements one the SpectralInfo object (labeled "info"), and the other the target dmd file (as a character labeled "dmdfile"). For <code>get_profile_all</code> it is a character vector with the *.dmd files that you want to retrieve.
...	other parameters for <code>plot.raster</code> .
dst_cluster	The spectra of the pixels that match which cluster/substance should be retrieved?
fpa	The fpa size, used in <code>get_profile_all</code>
waves	The wavenumbers, used in <code>get_profile_all</code>
plotpol	If true, the function will plot x and overlay contour lines highlighting <code>dst_cluster</code> . TRUE/FALSE.
plotpt	If true, the function will plot x and draw points over the pixels that match <code>dst_cluster</code> . For mosaics, the points will be only in the chunk selected in where. TRUE/FALSE.
cluster	If <code>class(x) == "SAM"</code> , Should the substances or the clusters be used for analysis? TRUE means that the clusters are used, FALSE that the substances. If you set the argument to FALSE, <code>dst_cluster</code> and <code>clusternames</code> should be adjusted. Use the name of the substance and the substances list from the SpectralReference .
slice	If the object is of class sam or Smooth , Which slice of them should be considered when retrieving the pixel locations?
clusternames	if <code>is.character(dst_cluster)</code> you should provide a vector with the clusternames to coerce <code>dst_cluster</code> to numeric.

Value

matrix with the columns matching the wavenumbers of 'where'.

Examples

```

x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
mosaic_sam(x, primpke, n_cores = 1)

```

```
y <- mosaic_compose(x@path, primpke@clusterlist)
y <- get_profile_sinfo(y,
  where = list("info" = x,
              "dmdfile" = "mosaic_0000_0000.dmd"),
  5, FALSE, FALSE)
```

highlight_substance *Highlight a selected substance*

Description

The function highlights a selected substance(s) -or cluster(s)- in a plot. It can add to an existing plot or create a new one. Currently implemented only for S3 objects returned by [clipper](#).

Usage

```
highlight_substance(
  x,
  dst_cluster,
  new = TRUE,
  clusternames = NULL,
  polygon_color = "red",
  match_uFTIR = FALSE,
  ...
)
```

Arguments

x	"clipper" object to plot
dst_cluster	cluster (or substance) to highlight. Numeric and character inputs are supported. If is char, then you should provide clusternames to coerce it to numeric.
new	Do you want a new plot or the function should add the polygons to the current plot?
clusternames	char vector given the list of analyzed substance clusters. Only relevant when dst_cluster is a character.
polygon_color	color of the overplotted polygons. Red by default.
match_uFTIR	The Agilent Microscope transposes and inverts the image (it inverts rows only - cols after transposition). Do you want the plot function to match this behaviour? Default FALSE.
...	other arguments for plot_tile

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
x <- tile_sam(x)
x <- smooth_sam(x, as.integer(length(primpke@clusternames)), window = 3, 1)
clip <- toClip(8,20,c(10,10))
plot(x)
polygon(clip@xycoords)
x <- clipper(x, clip@centre, clip@rad, 1)

highlight_substance(x, 5)
```

mask_sam

Mask SAM

Description

The spectral angle mapper algorithm always find a match, as it was designed to compare between to spectra. However, somethimes the match is far from good and the user could want to drop the poorly matched pixels. This function will create a mask for such a purpose.

Usage

```
mask_sam(x, threshold = 1.1)
```

Arguments

x an array. Typically, the array at the raw_sam slot of a SAM object.
threshold cut off value above wich, the SAM relation found is deemed insignificant

Value

S3 matrix

Examples

```
set.seed(4356)
x <- array(abs(rnorm(1000)), c(10, 10, 10))
x <- mask_sam(x, 0.1)
# trick to plot as clipper
class(x) <- c("clipper", "matrix")
plot(x, legend = FALSE)
```

matrix_sam	<i>Perform SAM between to matrices</i>
------------	--

Description

Perform the spectral angle mapper algorithm between to matrices.

Usage

```
matrix_sam(x, y, x.wl, y.wl)
```

Arguments

x	matrix
y	matrix
x.wl	wavenumbers for x
y.wl	wavenumbers for y

Value

vector

mosaic_chunk	<i>Mosaic chunk</i>
--------------	---------------------

Description

The function read a single mosaic tile (.dmd file extension) using a [SpectralInfo](#) object as a guide to find the file. It returns an object of class [Tile](#), which can be (pre/post)processed as if it were a single tile.

Usage

```
mosaic_chunk(info, path, dmdfile, fpa, wl)
```

Arguments

info	Optional. SpectralInfo object.
path	Optional. Path to dmdfile. Overwrites the path on the info object.
dmdfile	Target *.dmd file to read.
fpa	If info is missing, you should indicate the fpa size of the chunk.
wl	If info is missing, you should indicate the wavenumbers at which you read your data

Value

A [Tile](#) object.

Examples

```
x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
y <- mosaic_chunk(info = x, dmdfile = "mosaic_0000_0000.dmd")
class(y)
```

mosaic_compose

Mosaic compose

Description

There are two ways to process a mosaic. One way -which should be the standard- is to do it internally in R by calling first [mosaic_info](#) and later [mosaic_sam](#).

The function will write in the path specified in the [SpectralInfo](#) object returned by [mosaic_info](#) a series of binary files holding the SAM results. In other words, the function does not load the results to R automatically. You can call this function to load them back to R in a single object, which will be of class [SAM](#).

Usage

```
mosaic_compose(  
  path = ".",  
  clusterlist = NULL,  
  nslices = NULL,  
  drop_raw = FALSE  
)
```

Arguments

- | | |
|-------------|---|
| path | Where are the binary files? you can use the 'path' slot of the SpectralInfo instead of enter it manually. |
| clusterlist | The clusterlist vector that you passed along with the SpectralReference in the call to mosaic_sam . |
| nslices | If you deal with large mosaics, you might want to load only a few of the sam matches. This argument allows you to define up to which match you want to load to R. |
| drop_raw | If you are not interested in the angles you can set this argument to TRUE and avoid load them to R. |

Value

An object of class [SAM](#).

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

```
x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
mosaic_sam(x, primpke, n_cores = 1)
y <- mosaic_compose(x@path, primpke@clusterlist)
```

mosaic_info

Mosaic Info

Description

The function is equivalent to [tile_read](#). It makes an [SpectralInfo](#) object using the .dmt file. It does not load the measured spectra, and it was written for mosaic images only. See the page of [SpectralInfo](#) class for a full description of the returned object.

Usage

```
mosaic_info(dmtfile)
```

Arguments

dmtfile Path to the .dmt file.

Value

An object of class [SpectralInfo](#).

See Also

[SpectralInfo](#)

Examples

```
x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
```

mosaic_sam

*Mosaic SAM***Description**

The function performs the spectral angle mapper algorithm chunk by chunk for the mosaics files. It uses the parallel package, and by default all cores -1.

The function uses [mosaic_chunk](#) to cast a mosaic tile as a [Tile](#) object. Then performs a [tile_base_corr](#), [wavealing](#), and [tile_sam](#) to finally write binary files that hold the Spectral Angle Mapper results in the destination folder (path slot of the [SpectralInfo](#) object). The files can be loaded back to R using `link{mosaic_compose}`.

The function is using the parallel package, as each `tile_sam` takes a while to complete.

Usage

```
mosaic_sam(
  info,
  sref,
  derivative = NULL,
  base_corr = TRUE,
  FUN = NULL,
  n_cores = NULL
)
```

Arguments

<code>info</code>	SpectralInfo object.
<code>sref</code>	SpectralReference object.
<code>derivative</code>	whether to apply the first (1) or second (2) derivative before sam. Default NULL.
<code>base_corr</code>	TRUE/FALSE should tile_base_corr be call before processing each chunk?
<code>FUN</code>	A function to be passed to preprocess .It is always applied as if 'data' were a SpectralPack object.
<code>n_cores</code>	The number of cores to parallelize the task. NULL means all cores -1.

Value

TRUE

See Also

For a single tile application see [tile_sam](#).

Examples

```
x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
mosaic_sam(x, primpke, n_cores = 1)
```

plot,plot.uFTIR,missing-method
Plotting Objects

Description

Plotting method for objects of class [Tile](#), [SpectralPack](#), [SAM](#), [Smooth](#), and [clipper](#).

In objects of class SAM, always the clusters slot gets plotted.

Usage

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

plot_tile(x, slice = 1, FUN = sum, match_uFTIR = FALSE, ...)
```

Arguments

x	Tile, SpectralPack, SAM, Smooth, or clipper to plot.
y	Missing.
...	Further arguments to plot
slice	For objects of class SAM or Smooth, Which slice of the cube should be plotted?
FUN	For objects of class SpectralPack and Tile, Which function should be used to collapse the cube to a matrix?
match_uFTIR	The Agilent Microscope transposes and inverts the image (it inverts rows only - cols after transposition). Do you want the plot function to match this behaviour? Default FALSE.

Details

The function requires the raster package to plot. It coerces the matrix to raster to do so.

Examples

```
# Tile objects:
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
plot(x)
# with arguments for raster::plot
plot(x, axes = FALSE, box = FALSE, legend = FALSE)

# SpectralPack objects:
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
plot(x)

# SAM objects:
```

```

x <- tile_sam(x)
plot(x)

# Smooth objects
x <- smooth_sam(x, as.integer(length(primpke@clusternames)), window = 3, 1)
plot(x)

# clipper objects:
clip <- toClip(8,20,c(10,10))
polygon(clip@xycoords)
x <- clipper(x, clip@centre, clip@rad, 1)
plot(x)

```

preprocess

Tile Preprocess

Description

The function is a wrapper that allows the user to perform a user-define function to each pixel spectrum. It can be called internally in `mosaic_sam` using the FUN argument when processing mosaics.

As exemplified below, the function allows you to interact with other R packages that provide common features to analyze spectral data. Beware that you will need to adjust the wavenumbers manually if you are resampling... unless that 'data' is an object of class `SpectralPack`.

Usage

```

preprocess(data, FUN)

## S4 method for signature 'Tile,`function`'
preprocess(data, FUN)

## S4 method for signature 'SpectralPack,`function`'
preprocess(data, FUN)

```

Arguments

data	An object of class <code>Tile</code> or <code>SpectralPack</code> .
FUN	A function to apply to the Spectra slot.

Value

The same object but with the Spectra slot updated by FUN. `SpectralPack` objects return the wavelengths adjusted to the new extent, numbered from 1 to n (original values are lost).

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- preprocess(x, function(x){x+1})

# The function allows interacting with other R packages that provide common features
# for spectra analysis. For example, you can use the package "signal" to run
# a Savitzky-Golay filter.

if(requireNamespace("signal", quietly = TRUE)){

# for Tile objects. NOTE that after the preprocess x@wavenumbers does not match dim(x@Spectra)[3]
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- preprocess(x, function(x){signal::sgolayfilt(x)})
dim(x@Spectra)[3] == length(x@wavenumbers) # BEWARE!

# for SpectralPack objects
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
preprocess(x, function(x){signal::sgolayfilt(x)})

# Here the problem with the wavenumbers is gone
dim(x@Readings@Spectra)[3] == length(x@Readings@wavenumbers)
dim(x@Reference@Spectra)[3] == length(x@Reference@wavenumbers)
length(x@Readings@wavenumbers) == length(x@Reference@wavenumbers)

}
```

primpke

Example library

Description

Example spectral library to be used in microplastic evaluations. An example on how to format a [SpectralReference](#) object.

It was taken form:

Primpke, S., Wirth, M., Lorenz, C., Gerds, G. 2018. Reference database design for the automated analysis of microplastic samples based on Fourier transform infrared (FTIR) spectroscopy. *Analytical and Bioanalytical Chemistry* 410: 5131-5141. You might access to the article here <https://doi.org/10.1007/s00216-018-1156-x>

Usage

```
primpke
```

Format

an S4 `SpectralReference` object with:

substances character vector with the 270 names of the polymers included in Spectra

clusterlist integer vector with the corresponding cluster of each polymer described

clusternames the name of the cluster, sorted in ascending order according to clusterlist

Spectra matrix with the absorbance readings of each polymer (270 x 1176)

wavenumbers read wavenumbers. Correspond to Spectra ncol

Source

<https://doi.org/10.1007/s00216-018-1156-x>

rgdal_polygonize	<i>GDALPolygonize in R</i>
------------------	----------------------------

Description

The function calls the C GDALPolygonize routine (using C++). It does overwrite. It uses always 8 point connectedness. It writes on disc a file in "ESRI Shapefile" format.

I wrote a separate package with the function at first. However, I did not want to submit such a small package to CRAN to resolve the 'polygonize' problem. Therefor, I included the source code of that package in uFTIR at a later stage. If you are looking for a package to polygonize a raster file (similar to GDAL polygonize.py) you can use the GDALPolygonize package directly. You can find it in [GitHub](#).

That's also why this function is not exported for you to use it.

Usage

```
rgdal_polygonize(raster, folder, layer, field, overwrite = TRUE)
```

Arguments

raster	name/location of raster file to polygonize. Any driver included in GDAL should work.
folder	name/location of the shape file to write. File extension should not be included.
layer	name of the layer (.shp file) to write. Do not include de file extension.
field	name of the field that will keep the raster values.
overwrite	TRUE/FALSE to overwrite an existing file (folder).

Value

integer 0

Examples

```
NULL
```

SAM-class

Sam Prediction

Description

The class holds the results of the Spectral Angle Mapping algorithm and it originates by a call to [tile_sam](#) or to [mosaic_compose](#). It has three slots with congruent cubes. Each cube has its first two dimensions equal to the FPA size of the uFTIR Microscope image processed and the third dimension equal to the number of substances in the [SpectralReference](#) passed when calling [tile_sam](#) or [mosaic_sam](#). This class have methods to plot, summarize, and extract the readings of pixel matching a given [SpectralReference](#) substance. You can use the generics [plot](#), [summary](#), and [get_profile](#) but beware, you might have to provide extra arguments! (so, please don't see only the generic but the method for this class, see the section see also)

The [sam](#) object can be postprocessed by calling [smooth_sam](#) and/or clipped to a given extent by calling [clipper](#) (currently only circular extents are supported).

Value

An S4 object of class SAM

Slots

[raw_sam](#) a cube -array-. The numeric score for each [SpectralReference](#) substance passed by the call (slices), for each pixel of the FPA array of the [Tile](#) (rows and cols).

[substances](#) a cube -array-. The slices of the cube hold the substances (as number) of the [SpectralReference](#) object passed when calling [tile_sam](#) or [mosaic_sam](#). The substance number are sorted in a decreasing order. In this way, the first slide is the best match, the second the second-best, and so on according to the Spectral Angle Mapping algorithm.

[clusters](#) a cube -array-. Equivalent to the cube in the substances slot but instead of a substance number it has per ech slice the cluster number that corresponds to the clusterlist slot of the [SpectralReference](#) object.

See Also

[plot_tile](#) [summary_sam](#) [get_profile_tile](#) [get_profile_sinfo](#)

Examples

NULL

sam_load	<i>Sam load</i>
----------	-----------------

Description

The inverse function for sam_write

Usage

```
sam_load(dmdfile)
```

Arguments

dmdfile a connection to the .bin or .ban file (yes, is true).

Value

An object of [SAM](#)

sam_write	<i>Write a sam object to disk</i>
-----------	-----------------------------------

Description

A wrapper to the C++ mosaic_sam_write

Usage

```
sam_write(tile, sam)
```

Arguments

tile The tile form which the sam originates
sam The sam that you want to write to disk

Smooth-class	<i>Smooth</i>
--------------	---------------

Description

An object of this class is returned by a call to [smooth_sam](#). It holds a smoother version of a SAM object, keeping only the clusters slot and slicing it to a user-given number of slices. This class have methods to plot, summarize, and extract the readings of pixel matching a given SpectralReference substance. You can use the generics [plot](#), [summary](#), and [get_profile](#) but beware, you might have to provide extra arguments! (so, please don't see only the generic but the method for this class, see the section see also).

Value

An S4 object of class "Smooth".

Slots

`smooth` a cube -array- (even if only has 1 slice) that holds a smooth version of a SAM clusters slot.

See Also

[plot_tile](#) [summary_sam](#) [get_profile_tile](#) [get_profile_sinfo](#)

Examples

NULL

<code>smooth_sam</code>	<i>Smooth a Spectral Angle Mapper output</i>
-------------------------	--

Description

The function makes a smooth version of the object returned by [tile_sam](#) or [mosaic_compose](#). The function smooths out only the cluster slot, slicing it to a user-given range of slices.

Usage

```
smooth_sam(x, nclusters, window = 5, nslices = 1)
```

Arguments

x	Object of class SAM .
nclusters	How many clusters did the spectral library had? This parameter refer back to the length of the clusterlist slot of the SpectralReference object used when calling tile_sam or mosaic_sam . It should be an integer.
window	The function smooths out the SAM objectc "x" using a moving window. You should decide which size (in pixels) this window should be.
nslices	Integer. Starting from 1, up to which slice of the cube holded in the clusters slot of the SAM objectc "x" do you want the smooth_sam function to work? It will influence the output. See return.

Value

An S4 object of class [Smooth](#). It has only one slot (smooth) that holds an array in which each slice is the smooth version of the corresponding slice of a [SAM](#) object.

The returned object has methods to plot, summarize, and extract profiles by calling the generics [plot](#), [summary](#), and [get_profile](#). You might have to provide extra arguments. You can refer to the see also section to look for the object specific methods.

See Also

[plot_tile](#) [summary_sam](#) [get_profile_tile](#) [get_profile_sinfo](#)

Examples

```
a <- matrix(c(1,1,1,1,3,3,
              2,2,2,1,2,3,
              3,1,1,2,3,1,
              3,3,2,2,1,1,
              1,1,1,3,3,2,
              3,3,3,2,2,3), nrow = 6, byrow = TRUE)
a <- array(c(a,a), dim = c(6,6,2))
test_sam <- new(Class = "SAM",
               raw_sam = a,
               substances = a,
               clusters = a)

smooth_sam(test_sam, nclusters = 3, window = 3)
```

Description

General class that holds basic information about an image. Usually used when working with mosaics, as in this case the image is not read/loaded to R until is processed (e.g. by running a `mosaic_sam` process). Currently, the only function that returns an object of this class is he `mosaic_info` function. The object can by used later to read/load to R a single mosaic chunk using the `mosaic_chunk` function, or to process the whole mosaic image by calling the `mosaic_sam` function.

The class `Tile` contains the "Spectral_info" class. By this means, an object of class `Tile` holds the same information about the reading, and an extra slot called "Spectra" that holds the actual readings.

Value

An S4 object of class "Spectral_info"

Slots

`file` character. The full path to the .dmt or .bsp file.

`date` POSIXct. The date in which the image (mosaic or tile) was taken.

`fpsize` numeric. The size of the focal plane array used when measuring.

`wavenumbers` numeric. The wavenumbers at which the tarjet image (mosaic or tile) was taken.

`path` character. The path to the folder that holds all the uFTIR Microscope output files.

See Also

[Tile](#)

Examples

NULL

SpectralPack-class *Spectral Pack*

Description

A call to the function `wavealign` (specifically a call to `TileRead.wavealign`) returns an object of class `SpectralPack`. This class reunites the uFTIR Microscope readings loaded to R by a call to `tile_read` with a tarjet spectral library of class `SpectralReference`. Since this object is an output of a call to `wavealign` both the `Tile` object and `SpectralReference` have congruent wavenumbers. See `wavealign` how the resampling is done when calling different methods. A `SpectralPack` object is used as imput when calling `tile_sam` to run the Spectral Angle Mapping algorithm.

Value

An S4 object of class "SpectralPack" ready for spectral angle mapping.

Slots

Readings an object of class `Tile`.

Reference an object of class `SpectralReference`.

Examples

NULL

SpectralReference-class

Spectral Reference Class

Description

I made this class to hold spectral libraries. This libraries can be used to match a know spectrum with a pixel reading. `SpectralReference` objects are used by several functions in the package, at all analysis stages (i.e. preprocessing, processing, and summarizing/plotting). You can see an example in the data `primpke` (you can load it by calling `data("primpke")`). The format includes character/integer vectors to hold the names of the polymers, cluster IDs, and cluster names, all of which should correspond to one another. It also hold a matrix with the spectra of each polymer, with rows matching the vectors provided and a fourth vector holding the wavelenghts (that should be equal to the matrix cols).

The example provided in the package was taken form Primpke, S., Wirth, M., Lorenz, C., Gerdts, G. 2018. Reference database design for the automated analysis of microplastic samples based on Fourier transform infrared (FTIR) spectroscopy. *Analytical and Bioanalytical Chemistry* 410: 5131-5141. You might access to the article here <https://doi.org/10.1007/s00216-018-1156-x>.

If you have reference data that was taken considering a different range of wavenumbers, you should resample it first for the wavenumbers to match.

Value

A S4 object of class "SpectralReference".

Slots

`substances` A character vector identifying by name the polymers whose spectrum is provided in `Spectra`. The order of the elements should be consistent with `clusterlist`, `clusternames` and `Spectra` rows. Along the same lines, the object expects `length(substances) == nrow(Spectra)`.

`clusterlist` If the polymers are aggregated in clusters this slot should hold an integer vector with the correspondent cluster for each polymer individuated in `substances`. If you don't want to use clusters or you don't have clusters for your polymers/library you can place a sequence of integers from `1:nrow(Spectra)` (e.g. by calling `seq_along(substances)`) in this slot.

`clusternames` Character vector holding reference names for each cluster, sorted according to `clusterlist`. In other words, if you want to name cluster 1L "polystyrene" the first position in `clusternames` should be equal to "polystyrene". If you don't want to use clusters or you don't have clusters for your polymers/library you can duplicate `substances`. The program expects that the length of the `clusternames` vector equals the length of unique elements in `clusterlist`.

Spectra A matrix holding row-wise the spectrum of each substance. Each row should correspond to the spectrum of one substance. In the same lines, columns should hold recorded measures for each wavenumber. The program expects that the length of the substance vector equals the number of rows of Spectra and that the number of columns of Spectra equals the length of the wavenumbers vector.

wavenumbers A numeric vector with the wavenumbers.

Examples

```
data("primpke")
```

```
summary,summary_sam-method
```

Method to summarize SAM or Smooth objects.

Description

Summary method for SAM and Smooth objects.

Usage

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

```
summary_sam(
  object,
  mask = NULL,
  clusternames = NULL,
  slice = 1,
  window = NULL,
  smooth = TRUE,
  temporal = FALSE
)
```

Arguments

<code>object</code>	An object of class SAM or Smooth . It accepts also object of S3 class "clipper" returned by the clipper function.
<code>...</code>	other parameters for <code>summary_sam</code>
<code>mask</code>	a clipper mask returned by toClip function. It should be set to NULL To avoid clipping, the default.
<code>clusternames</code>	The name of the clusters, it should match the clusternames list comprised in the SpectralReference object passed to tile_sam or mosaic_sam . Mandatory if object is of class SAM when <code>smooth = TRUE</code> .
<code>slice</code>	Which slice of object should be summarized?
<code>window</code>	When <code>smooth = TRUE</code> the size of the window for smooth_sam .

smooth	When object if of class <code>SAM</code> , should the function silently apply <code>smooth_sam</code> before summarizing?.
temporal	if TRUE the raster and shape files from which the summary is extracted are written in the working directory.

Value

A dataframe with three columns showing the cluster number, the area in pixels², and the cluster name as character (only if the parameter `clusternames` is within the function call). Each row stand for a single particle. The function writes on disk two files "raster_out.tif" which is a raster of the object passed through the object parameter, and a shape_out "ESRI Shapefile" folder with a shapefile holding a vectorized version of "raster_out.tif".

Examples

```
x <- mosaic_info(base::system.file("extdata/mosaic.dmt", package = "uFTIR"))
mosaic_sam(x, primpke, n_cores = 1)
y <- mosaic_compose(x@path, clusterlist = primpke@clusterlist)
summary_sam(y, clusternames = primpke@clusternames, smooth = FALSE, temporal = TRUE)
```

Tile-class

Tile

Description

Class to hold the reading of a single tile. It can also store a single tile of a mosaic, to which we will refer to as chunks. It is an extended class for `SpectralInfo`, adding an extra slot to hold the uFTIR Microscope readings -called Spectra. The functions `tile_read` and `mosaic_chunk` return an object of this class. The class has a defined method to plot, which can be called either by the generic `plot` or by `plot_tile`. You can use an object of this class in the following functions:

`plot_tile` To plot a heat map of the reading. Available too through the generic `plot`.

`tile_base_corr` A preprocessing function that replace negative values with zeros.

`wavealign` A resampling method that allows to match the wavenumbers of a given tile/chunk with the wavenumbers of a `SpectralReference`.

`tile_sam` When a Tile object is linked to a `SpectralReference` by a former call to `wavealign`, this function performs a spectral angle mapper match between the Tile and the `SpectralReference`.

`get_profile_tile` To get the pixels spectra that matched a given polymer/cluster.

Value

An S4 object of class "Tile".

Slots

`Spectra` array. The aquired spectra (.dat or dmd file)

Examples

```
NULL
```

tile_base_corr	<i>Baseline correction</i>
----------------	----------------------------

Description

A function to subtract the minimum of a pixel (vector) to have a minimal signal = 0, avoiding negative readings. Currently the only preprocessing method implemented.

Usage

```
tile_base_corr(data)
```

Arguments

data Object of class [Tile](#).

Value

An object of class [Tile](#) congruent with data, with the spectra modified.

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
```

tile_read	<i>Single tile read</i>
-----------	-------------------------

Description

The function loads a single tile to R as an object of class [Tile](#).

The function is also capable to load the toy mosaic produced by the Agilent software. You can do so by passing as bspfile parameter the .bsp file comprised in a mosaic folder. However, beware! the toy mosaic IS NOT the complete mosaic, it is only a reduced version of it (i.e. you will lose resolution, and you will not longer know the pixel size). Use it with care.

Usage

```
tile_read(bspfile)
```

Arguments

bspfile path to the .bsp file.

Value

An S4 Object of class [Tile](#).

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
```

tile_sam

Spectral Angle Mapper

Description

Performs the Spectral Angle Mapper to match the [SpectralReference](#) with your readings (as [Tile](#)).

Usage

```
tile_sam(SpectralPack, derivative = NULL)
```

Arguments

SpectralPack an object of class [SpectralPack](#).
derivative whether to apply the first (1) or second (2) derivative to preprocess the data before the algorithm is applied. Default NULL (It tells the program to not derivate).

Value

An object of class [SAM](#).

See Also

For its application to mosaic images see [mosaic_sam](#).

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))  
x <- tile_base_corr(x)  
x <- wavealign(x, primpke)  
x <- tile_sam(x)
```

`toClip`*Visual clipper*

Description

Visual aid to find the right center and radius for the `clipper` function. It does not clip, but return an usable (plotable) object.

Since the samples are placed under the uFTIR Microscope by hand, the cropping area to pass to `clipper` is not always the same and (usually) it has to be adjusted. To have a visual aid, you can use this function. It returns an S4 object of class `clipmask` which can be over-plotted on top of a `SAM` or `Smooth` object by calling `polygon` and using the `xycoords` slot of the returned object. Using that process you can test manually different center points and radius for the clipping circle.

Usage

```
toClip(rad = 1, segments = 5, centre = c(0, 0))
```

Arguments

<code>rad</code>	The circle radius.
<code>segments</code>	How many segments should the resulting polygon have? (because it is not a circle, 20 is ok).
<code>centre</code>	The coordinates of the polygon centre (x,y).

Value

An object of class `clipmask`.

See Also

`clipper`

Examples

```
toClip(1, 5, c(0,0))

x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
x <- tile_sam(x)
x <- smooth_sam(x, as.integer(length(primpke@clusternames)), window = 3, 1)

clip <- toClip(8,20,c(10,10))
polygon(clip@xycoords)

x <- clipper(x, clip@centre, clip@rad, 1)
```

uFTIR

*Process and analyze Agilent Cary 620 FTIR Microscope images***Description**

Process and analyze Agilent Cary 620 FTIR Microscope images

wavealign

*Methods to resample the wavenumbers***Description**

It clips and resamples the wavenumbers of `data.x` and `data.y` to a common extent. Typically, it uses the wavenumbers of `data.x` to resample `data.y` and then the function clips the largest base on the narrowest.

Usage

```

wavealign(data.x, data.y)

## S4 method for signature 'numeric,SpectralReference'
wavealign(data.x, data.y)

## S4 method for signature 'Tile,SpectralReference'
wavealign(data.x, data.y)

## S4 method for signature 'SpectralReference,Tile'
wavealign(data.x, data.y)

MosaicGetRef.wavealign(data.x, data.y)

TileRead.wavealign(data.x, data.y)

```

Arguments

`data.x` An object of class [SpectralReference](#) or [Tile](#).
`data.y` An object of class [SpectralReference](#) or [Tile](#). It should be the other one.

Details

There are two methods defined:

- **TileRead.wavealign** - This method should be the only one used by the user.
- **MosaicGetRef.wavealign** - This method is called internally by `mosaic_sam`. The method is different since it expect a numeric vector in `data.x`, which is taken from the wavenumbers slot of a [SpectralInfo](#) object. The implication is that, for mosaics, the [SpectralReference](#) is always resampled according to the wavenumbers of the mosaics. It was programed this way, since mosaics are not loaded into the memory (R) until they are processed by `mosaic_sam`.

Value

- **TileRead.wavealign** - An object of class [SpectralPack](#)
- **MosaicGetRef.wavealign** - The [SpectralReference](#) object passed as data.y, resampled to data.x.

Examples

```
x <- tile_read(base::system.file("extdata/tile.bsp", package = "uFTIR"))
x <- tile_base_corr(x)
x <- wavealign(x, primpke)
```

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