

Package ‘OpenSpecy’

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Type Package

Title Analyze, Process, Identify, and Share Raman and (FT)IR Spectra

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Description Raman and (FT)IR spectral analysis tool for plastic particles and other environmental samples (Cowger et al. 2021, <[doi:10.1021/acs.analchem.1c00123](https://doi.org/10.1021/acs.analchem.1c00123)>). Supported features include reading spectral data files (.asp, .csv, .jdx, .spc, .spa, .0), Savitzky-Golay smoothing of spectral intensities with `smooth_intens()`, correcting background noise with `subtr_bg()` in accordance with Zhao et al. (2007) <[doi:10.1366/000370207782597003](https://doi.org/10.1366/000370207782597003)>, and identifying spectra using an onboard reference library (Cowger et al. 2020, <[doi:10.1177/0003702820929064](https://doi.org/10.1177/0003702820929064)>). Analyzed spectra can be shared with the Open Specy community. A Shiny app is available via `run_app()` or online at <<https://openanalysis.org/openspecy/>>.

URL <https://github.com/wincowgerDEV/OpenSpecy-package/>

BugReports <https://github.com/wincowgerDEV/OpenSpecy-package/issues/>

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adj_intens	<i>Adjust spectral intensities to absorbance units</i>
------------	--

Description

Converts reflectance or transmittance intensity units to absorbance units.

Usage

```
adj_intens(x, ...)

## S3 method for class 'formula'
adj_intens(formula, data = NULL, ...)
```

```
## S3 method for class 'data.frame'  
adj_intens(x, ...)  
  
## Default S3 method:  
adj_intens(x, y, type = "none", make_rel = TRUE, ...)
```

Arguments

x	a numeric vector containing the spectral wavenumbers; alternatively a data frame containing spectral data as "wavenumber" and "intensity" can be supplied.
formula	an object of class 'formula' of the form intensity ~ wavenumber.
data	a data frame containing the variables in formula.
y	a numeric vector containing the spectral intensities.
type	a character string specifying whether the input spectrum is in absorbance units ("none", default) or needs additional conversion from "reflectance" or "transmittance" data.
make_rel	logical; if TRUE spectra are automatically normalized with make_rel() .
...	further arguments passed to the submethods.

Details

Many of the Open Specy functions will assume that the spectrum is in absorbance units. For example, see [match_spec\(\)](#) and [subtr_bg\(\)](#). To run those functions properly, you will need to first convert any spectra from transmittance or reflectance to absorbance using this function. The transmittance adjustment uses the $\log_{10}(1/T)$ calculation which does not correct for system and particle characteristics. The reflectance adjustment uses the Kubelka-Munk equation $(1 - R)^2/2R$. We assume that the reflectance intensity is a percent from 1-100 and first correct the intensity by dividing by 100 so that it fits the form expected by the equation.

Value

[adj_intens\(\)](#) returns a data frame containing two columns named "wavenumber" and "intensity".

Author(s)

Win Cowger, Zacharias Steinmetz

See Also

[subtr_bg\(\)](#) for spectral background correction; [match_spec\(\)](#) matches spectra with the Open Specy or other reference libraries

Examples

```
data("raman_hdpe")  
  
adj_intens(raman_hdpe)
```

`adj_neg`*Normalization of spectral data*

Description

`adj_neg()` converts numeric values $x < 1$ into values ≥ 1 , keeping absolute differences between values by shifting intensity values with the value of the smallest number. `make_rel()` converts values x into relative values between 0 and 1 using the standard normalization equation. If `na.rm` is TRUE, missing values are removed before the computation proceeds.

Usage

```
adj_neg(x, na.rm = FALSE)
```

```
make_rel(x, na.rm = FALSE)
```

Arguments

<code>x</code>	a numeric vector or an R object which is coercible to one by <code>as.vector(x, "numeric")</code> ; <code>x</code> should be intensity data.
<code>na.rm</code>	logical. Should missing values be removed?

Details

`adj_neg()` is used in Open Specy to avoid errors that could arise from log transforming spectra when using `adj_intens()` and other functions. `make_rel()` is used in Open Specy to retain the relative height proportions between spectra while avoiding the large numbers that can result from some spectral instruments.

Value

`adj_neg()` and `make_rel()` return numeric vectors with the normalized data.

Author(s)

Win Cowger, Zacharias Steinmetz

See Also

`min()` for the calculation of minima; `adj_intens()` for log transformation functions

Examples

```
adj_neg(c(-1000, -1, 0, 1, 10))  
make_rel(c(-1000, -1, 0, 1, 10))
```

Description

These functions will import the spectral libraries from Open Specy if they were not already downloaded. The CRAN does not allow for deployment of large datasets so this was a workaround that we are using to make sure everyone can easily get Open Specy functionality running on their desktop.

Usage

```
check_lib(  
  which = c("ftir", "raman"),  
  types = c("metadata", "library", "peaks"),  
  path = "system",  
  condition = "warning"  
)  
  
get_lib(  
  which = c("ftir", "raman"),  
  types = c("metadata", "library", "peaks"),  
  path = "system",  
  node = "x7dpz",  
  conflicts = "overwrite",  
  ...  
)  
  
load_lib(  
  which = c("ftir", "raman"),  
  types = c("metadata", "library", "peaks"),  
  path = "system"  
)
```

Arguments

which	a character string specifying which library to use, "raman" or "ftir".
types	library types to check/retrieve; defaults to c("metadata", "library", "peaks").
path	where to save or look for local library files; defaults to "system" pointing to <code>system.file("extdata", package = "OpenSpecy")</code> .
condition	determines if <code>check_lib()</code> should warn ("warning", the default) or throw and error ("error").
node	the OSF node to be retrieved; should be "x7dpz".
conflicts	determines what happens when a file with the same name exists at the specified destination. Can be one of the following (see <code>osf_download()</code> for details):

- "error" throw an error and abort the file transfer operation.
- "skip" skip the conflicting file(s) and continue transferring the remaining files.
- "overwrite" (default) replace the existing file with the transferred copy.

... further arguments passed to `osf_download()`.

Details

`check_lib()` checks to see if the Open Specy reference library already exists on the users computer. `get_lib()` downloads the Open Specy library from OSF ([doi:10.17605/OSF.IO/X7DPZ](https://doi.org/10.17605/OSF.IO/X7DPZ)). `load_lib()` will load the library into the global environment for use with the Open Specy functions.

Value

`check_lib()` and `get_lib()` return messages only; `load_lib()` returns a list object containing the respective spectral reference library.

Author(s)

Zacharias Steinmetz

References

Cowger W, Gray A, Christiansen SH, Christiansen SH, Christiansen SH, De Frond H, Deshpande AD, Hemabessiere L, Lee E, Mill L, et al. (2020). "Critical Review of Processing and Classification Techniques for Images and Spectra in Microplastic Research." *Applied Spectroscopy*, **74**(9), 989–1010. [doi:10.1177/0003702820929064](https://doi.org/10.1177/0003702820929064).

Cowger, W (2021). "Library data." *OSF*. [doi:10.17605/OSF.IO/X7DPZ](https://doi.org/10.17605/OSF.IO/X7DPZ).

See Also

[match_spec\(\)](#)

Examples

```
## Not run:
check_lib(which = c("ftir", "raman"))
get_lib(which = c("ftir", "raman"))

spec_lib <- load_lib(which = c("ftir", "raman"))

## End(Not run)
```

human_ts	<i>Create human readable timestamps</i>
----------	---

Description

This helper function creates human readable timestamps in the form of %Y%m%d-%H%M%OS at the current time.

Usage

```
human_ts()
```

Details

Human readable timestamps are appended to file names and fields when metadata are shared with the Open Specy community.

Value

human_ts() returns a character value with the respective timestamp.

Author(s)

Win Cowger

See Also

[format.Date](#) for date conversion functions

Examples

```
human_ts()
```

match_spec	<i>Match spectra with reference library</i>
------------	---

Description

match_spec() will compare a spectrum to a spectral library formatted with the Open Specy standard and report the best match using the Pearson correlation coefficient. find_spec() makes it easy to retrieve single spectra and metadata from the Open Specy reference library.

Usage

```

match_spec(x, ...)

## S3 method for class 'formula'
match_spec(formula, data = NULL, ...)

## S3 method for class 'data.frame'
match_spec(x, ...)

## Default S3 method:
match_spec(
  x,
  y,
  library,
  which = NULL,
  type = "full",
  range = seq(0, 6000, 0.1),
  top_n = 100,
  ...
)

find_spec(
  subset,
  library,
  which = NULL,
  type = "metadata",
  cols = c("spectrum_identity", "organization", "contact_info", "spectrum_type",
    "instrument_used", "instrument_accessories", "instrument_mode", "laser_light_used",
    "total_acquisition_time_s", "number_of_accumulations",
    "level_of_confidence_in_identification", "cas_number", "material_producer",
    "material_purity", "material_form", "material_quality", "spectral_resolution",
    "data_processing_procedure", "other_information", "sample_name", "wavenumber",
    "intensity", "group"),
  ...
)

```

Arguments

x	a numeric vector containing the spectral wavenumbers; alternatively a data frame containing spectral data as "wavenumber" and "intensity" can be supplied.
formula	an object of class 'formula' of the form intensity ~ wavenumber.
data	a data frame containing the variables in formula.
y	a numeric vector containing the spectral intensities.
library	reference library you want to compare against.
which	a character string specifying which library to match, "raman" or "ftir".

type	a character string specifying whether the "full" spectrum should be matched or spectrum "peaks" only. "metadata" is needed to browser spectra with <code>find_spec()</code> .
range	this should be all possible wavenumber values from your spectral library.
top_n	number of top matches that you want to be returned.
subset	logical expression indicating elements or rows to search for; see <code>subset()</code> for details.
cols	columns to retrieve from the Open Specy reference library; columns containing no or missing values are automatically removed.
...	further arguments passed to the submethods.

Details

This routine will match the spectrum you want to identify to the wavenumbers present in the spectral library. Once the spectra are aligned, it computes the Pearson correlation coefficient between the spectrum you want to identify and all spectra in the library (see `cor`). The function returns a table with the Pearson correlation coefficient values and all metadata for the top spectral matches. If using the Open Specy library, all intensity values are in absorbance, so your spectra should also be in absorbance units. If you need to convert your spectrum, use `adj_intens()`.

Value

`match_spec()` returns a data frame with the `top_n` material matches, their Pearson's r value, and the organization they were provided by. `find_spec()` returns a data frame with the spectral raw data or metadata of a specific reference spectrum.

Author(s)

Win Cowger, Zacharias Steinmetz

See Also

`adj_intens()` converts spectra; `get_lib()` retrieves the Open Specy reference library; `load_lib()` loads the Open Specy reference library into an R object of choice

Examples

```
## Not run:
data("raman_hdpe")

get_lib("raman")
spec_lib <- load_lib("raman")

match_spec(raman_proc, library = spec_lib, which = "raman")

find_spec(sample_name == 5381, library = spec_lib, which = "raman")

## End(Not run)
```

raman_hdpe	<i>Sample Raman spectrum</i>
------------	------------------------------

Description

Raman spectrum of high-density polyethylene (HDPE).

Format

A data table containing 964 rows and 2 columns:

wavenumber:	spectral wavenumber [1/cm]
intensity:	absorbance values [-]

Author(s)

Win Cowger

References

Cowger W, Gray A, Christiansen SH, Christiansen SH, Christiansen SH, De Frond H, Deshpande AD, Hemabessiere L, Lee E, Mill L, et al. (2020). "Critical Review of Processing and Classification Techniques for Images and Spectra in Microplastic Research." *Applied Spectroscopy*, **74**(9), 989–1010. doi:10.1177/0003702820929064.

Examples

```
data("raman_hdpe")
```

read_text	<i>Read spectral data</i>
-----------	---------------------------

Description

Functions for reading spectral data types including .asp, .jdx, .spc, .spa, .0, and .csv.

Usage

```
read_text(  
  file = ".",  
  cols = NULL,  
  method = "read.csv",  
  share = NULL,  
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
```

```
    ...
  )

read_asp(
  file = ".",
  share = NULL,
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

read_spa(
  file = ".",
  share = NULL,
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

read_jdx(
  file = ".",
  share = NULL,
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

read_spc(
  file = ".",
  share = NULL,
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

read_0(
  file = ".",
  share = NULL,
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

read_extdata(file = NULL)
```

Arguments

file	file to be read from.
cols	character vector of length = 2 indicating the column names for the wavenumber and intensity; if NULL columns are guessed.
method	submethod to be used for reading text files; defaults to read.csv but fread works as well.

share	defaults to NULL; needed to share spectra with the Open Specy community; see share_spec() for details.
id	a unique user and/or session ID; defaults to <code>paste(digest(Sys.info()), digest(sessionInfo()), sep = "/")</code> .
...	further arguments passed to the submethods.

Details

`read_spc()` and `read_jdx()` are just a wrapper around the functions provided by the [hyperSpec](#) package. Other functions have been adapted various online sources. All functions convert datasets to a 2 column table with one column labeled "wavenumber" and the other "intensity". There are many unique iterations of spectral file formats so there may be bugs in the file conversion. Please contact us if you identify any.

Value

All `read_*`() functions return data frames containing two columns named "wavenumber" and "intensity".

Author(s)

Zacharias Steinmetz, Win Cowger

See Also

[read.jdx\(\)](#); [read.spc\(\)](#); [readRaw\(\)](#); [share_spec\(\)](#)

Examples

```
read_text(read_extdata("raman_hdpe.csv"))
read_asp(read_extdata("ftir_ldpe_soil.asp"))
read_0(read_extdata("ftir_ps.0"))
```

run_app

Run Open Specy app

Description

This wrapper function starts the graphical user interface of Open Specy.

Usage

```
run_app(app_dir = "system", path = "system", log = TRUE, ...)
```

Arguments

app_dir	the app to run; defaults to "system" pointing to <code>system.file("shiny", package = "OpenSpecy")</code> .
path	where to look for the local library files; defaults to "system" pointing to <code>system.file("extdata", package = "OpenSpecy")</code> .
log	logical; enables/disables logging to <code>tempdir()</code>
...	arguments passed to <code>runApp()</code> .

Details

After running this function the Open Specy GUI should open in a separate window or in your computer browser.

Value

This function normally does not return any value, see `runApp()`.

Author(s)

Zacharias Steinmetz

See Also

`runApp()`

Examples

```
## Not run:  
run_app()  
  
## End(Not run)
```

share_spec

Share data with the Open Specy community

Description

This helper function shares spectral data and metadata with the Open Specy community.

Please note that `share_spec()` only provides basic sharing functionality if used interactively. This means that files are only formatted and saved for sharing but are not send automatically. This only works with hosted instances of Open Specy.

Usage

```

share_spec(data, ...)

## Default S3 method:
share_spec(data, ...)

## S3 method for class 'data.frame'
share_spec(
  data,
  metadata = c(user_name = "", contact_info = "", organization = "", citation = "",
    spectrum_type = "", spectrum_identity = "", material_form = "", material_phase = "",
    material_producer = "", material_purity = "", material_quality = "", material_color =
      "", material_other = "", cas_number = "", instrument_used = "",
    instrument_accessories = "", instrument_mode = "", spectral_resolution = "",
    laser_light_used = "", number_of_accumulations = "", total_acquisition_time_s = "",
    data_processing_procedure = "", level_of_confidence_in_identification = "",

    other_info = "", license = "CC BY-NC"),
  file = NULL,
  share = "system",
  id = paste(digest(Sys.info()), digest(sessionInfo()), sep = "/"),
  ...
)

```

Arguments

data	a data frame containing the spectral data; columns should be named "wavenumber" and "intensity".
metadata	a named vector of the metadata to share; see details below.
file	file to share (optional).
share	accepts any local directory to save the spectrum for later sharing via e-mail to <wincowger@gmail.com>; "system" (default) uses the Open Specy package directory at <code>system.file("extdata", package = "OpenSpecy")</code> ; if a correct API token exists, "dropbox" shares the spectrum with the cloud.
id	a unique user and/or session ID; defaults to <code>paste(digest(Sys.info()), digest(sessionInfo()), sep = "/")</code> .
...	further arguments passed to the submethods.

Details

The metadata argument may contain a named vector with the following details (* = mandatory):

user_name*:	User name, e.g. "Win Cowger"
contact_info:	Contact information, e.g. "1-513-673-8956, wincowger@gmail.com"
organization:	Affiliation, e.g. "University of California, Riverside"
citation:	Data citation, e.g. "Primpke, S., Wirth, M., Lorenz, C., & Gerdt, G. (2018).
spectrum_type*:	Raman or FTIR

spectrum_identity*:	Material/polymer analyzed, e.g. "Polystyrene"
material_form:	Form of the material analyzed, e.g. textile fiber, rubber band, sphere, granule
material_phase:	Phase of the material analyzed (liquid, gas, solid)
material_producer:	Producer of the material analyzed, e.g. Dow
material_purity:	Purity of the material analyzed, e.g. 99.98%
material_quality:	Quality of the material analyzed, e.g. consumer product, manufacturer material
material_color:	Color of the material analyzed, e.g. blue, #0000ff, (0, 0, 255)
material_other:	Other material description, e.g. 5 µm diameter fibers, 1 mm spherical particles
cas_number:	CAS number, e.g. 9003-53-6
instrument_used:	Instrument used, e.g. Horiba LabRam
instrument_accessories:	Instrument accessories, e.g. Focal Plane Array, CCD
instrument_mode:	Instrument modes/settings, e.g. transmission, reflectance
spectral_resolution:	Spectral resolution, e.g. 4/cm
laser_light_used:	Wavelength of the laser/light used, e.g. 785 nm
number_of_accumulations:	Number of accumulations, e.g. 5
total_acquisition_time_s:	Total acquisition time (s), e.g. 10 s
data_processing_procedure:	Data processing procedure, e.g. spikefilter, baseline correction, none
level_of_confidence_in_identification:	Level of confidence in identification, e.g. 99%
other_info:	Other information
license:	The license of the shared spectrum; defaults to "CC BY-NC" (see https://creativecommons.org/licenses/by-nc/4.0/)

Value

share_spec() returns only messages/warnings.

Author(s)

Zacharias Steinmetz, Win Cowger

See Also

[read_text\(\)](#); [digest\(\)](#); [sessionInfo\(\)](#)

Examples

```
## Not run:
data("raman_hdpe")
share_spec(raman_hdpe,
           metadata = c(user_name = "Win Cowger",
                       spectrum_type = "FTIR",
                       spectrum_identity = "PE",
                       license = "CC BY-NC"),
           share = tempdir())

## End(Not run)
```

smooth_intens *Smooth spectral intensities*

Description

This smoother can enhance the signal to noise ratio of the data and uses a Savitzky-Golay filter with a running window of data points and the polynomial specified.

Usage

```
smooth_intens(x, ...)

## S3 method for class 'formula'
smooth_intens(formula, data = NULL, ...)

## S3 method for class 'data.frame'
smooth_intens(x, ...)

## Default S3 method:
smooth_intens(x, y, p = 3, n = 11, make_rel = TRUE, ...)
```

Arguments

x	a numeric vector containing the spectral wavenumbers; alternatively a data frame containing spectral data as "wavenumber" and "intensity" can be supplied.
formula	an object of class 'formula' of the form intensity ~ wavenumber.
data	a data frame containing the variables in formula.
y	a numeric vector containing the spectral intensities.
p	polynomial order for the filter
n	number of data points in the window, filter length (must be odd).
make_rel	logical; if TRUE spectra are automatically normalized with make_rel() .
...	further arguments passed to sgolay() .

Details

This is a wrapper around the filter function in the signal package to improve integration with other Open Specy functions. A typical good smooth can be achieved with 11 data point window and a 3rd or 4th order polynomial.

Value

smooth_intens() returns a data frame containing two columns named "wavenumber" and "intensity".

Author(s)

Win Cowger, Zacharias Steinmetz

References

Savitzky A, Golay MJ (1964). "Smoothing and Differentiation of Data by Simplified Least Squares Procedures." *Analytical Chemistry*, **36**(8), 1627–1639.

See Also

[sgolay\(\)](#)

Examples

```
data("raman_hdpe")
smooth_intens(raman_hdpe)
```

spec_res

Spectral resolution

Description

Helper function for calculating the spectral resolution from wavenumber data.

Usage

```
spec_res(x)
```

Arguments

x a numeric vector or an R object which is coercible to one by `as.vector(x, "numeric")`; x should be wavenumber data.

Details

The spectral resolution is the the minimum wavenumber, wavelength, or frequency difference between two lines in a spectrum that can still be distinguished.

Value

`spec_res()` returns a single numeric value.

Author(s)

Win Cowger, Zacharias Steinmetz

Examples

```
data("raman_hdpe")
spec_res(raman_hdpe$wavenumber)
```

subtr_bg

*Automated background subtraction for spectral data***Description**

This baseline correction routine iteratively finds the baseline of a spectrum using a polynomial fitting.

Usage

```
subtr_bg(x, ...)

## S3 method for class 'formula'
subtr_bg(formula, data = NULL, ...)

## S3 method for class 'data.frame'
subtr_bg(x, ...)

## Default S3 method:
subtr_bg(x, y, degree = 8, raw = FALSE, make_rel = TRUE, ...)
```

Arguments

x	a numeric vector containing the spectral wavenumbers; alternatively a data frame containing spectral data as "wavenumber" and "intensity" can be supplied.
formula	an object of class 'formula' of the form intensity ~ wavenumber.
data	a data frame containing the variables in formula.
y	a numeric vector containing the spectral intensities.
degree	the degree of the polynomial. Must be less than the number of unique points when raw is FALSE. Typically a good fit can be found with a 8th order polynomial.
raw	if TRUE, use raw and not orthogonal polynomials.
make_rel	logical; if TRUE spectra are automatically normalized with <code>make_rel()</code> .
...	further arguments passed to <code>poly()</code> .

Details

This is a translation of Michael Stephen Chen's MATLAB code written for the `imodpolyfit` routine from Zhao et al. 2007.

Value

`subtr_bg()` returns a data frame containing two columns named "wavenumber" and "intensity".

Author(s)

Win Cowger, Zacharias Steinmetz

References

Chen MS (2020). Michaelstchen/ModPolyFit. *MATLAB*. Retrieved from <https://github.com/michaelstchen/modPolyFit> (Original work published July 28, 2015)

Zhao J, Lui H, McLean DI, Zeng H (2007). "Automated Autofluorescence Background Subtraction Algorithm for Biomedical Raman Spectroscopy." *Applied Spectroscopy*, **61**(11), 1225–1232. doi:10.1366/000370207782597003.

See Also

`poly()`; `smooth_intens()`

Examples

```
data("raman_hdpe")  
  
subtr_bg(raman_hdpe)
```

test_lib

Test reference library

Description

Reference library of 34 Raman spectra used for internal testing.

Format

A list named "test" with two elements:

```
metadata:  metadata of 34 Raman spectra  
library:   all reference spectra, sample_name serves as identifier
```

Author(s)

Jennifer Lynch

Examples

```
data("test_lib")
```

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