

# Package ‘OpenSpecy’

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**Type** Package

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

**Version** 1.5.3

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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)>). With `read_any()`, Open Specy provides a single function for reading individual, batch, or map spectral data files like `.asp`, `.csv`, `.jdx`, `.spc`, `.spa`, `.0`, and `.zip`. `process_spec()` simplifies processing spectra, including smoothing, baseline correction, range restriction and flattening, intensity conversions, wavenumber alignment, and min-max normalization. Spectra can be identified in batch using an onboard reference library (Cowger et al. 2020, <[doi:10.1177/0003702820929064](https://doi.org/10.1177/0003702820929064)>)) using `match_spec()`. A Shiny app is available via `run_app()` or online at <<https://www.openanalysis.org/openspecy/>>.

**URL** <https://github.com/wincowgerDEV/OpenSpecy-package/>,  
<https://raw.githack.com/wincowgerDEV/OpenSpecy-package/main/docs/index.html>

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

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**Encoding** UTF-8

**LazyLoad** true

**LazyData** true

**VignetteBuilder** knitr

**Depends** R (>= 4.3.0)

**Imports** methods, data.table, jsonlite, yaml, caTools, hyperSpec, mmand, plotly, digest, signal, glmnet, jpeg, shiny

**Suggests** knitr, rmarkdown, testthat (>= 3.1.9), shinyjs, shinyWidgets, bs4Dash, dplyr, DT, ggplot2

**RoxygenNote** 7.3.2

**Config/testthat.edition** 3

**NeedsCompilation** no

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adj_intens	<i>Adjust spectral intensities to standard absorbance units.</i>
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**Description**

Converts reflectance or transmittance intensity units to absorbance units and adjust log or exp transformed units.

**Usage**

```
adj_intens(x, ...)

## Default S3 method:
adj_intens(x, type = "none", make_rel = TRUE, log_exp = "none", ...)

## S3 method for class 'OpenSpecy'
adj_intens(x, type = "none", make_rel = TRUE, log_exp = "none", ...)
```

**Arguments**

- |      |  |
|------|--|
| x    | a list object of class OpenSpecy.  |
| type | a character string specifying whether the input spectrum is in absorbance units ("none", default) or needs additional conversion from "reflectance" or "transmittance" data. |

<code>make_rel</code>	logical; if TRUE spectra are automatically normalized with <a href="#">make_rel()</a> .
<code>log_exp</code>	a character string specifying whether the input needs to be log transformed "log", exp transformed "exp", or not ("none", default).
...	further arguments passed to submethods; this is to <a href="#">adj_neg()</a> for adj_intens() and to <a href="#">conform_res()</a> for conform_intens().

## Details

Many of the Open Specy functions will assume that the spectrum is in absorbance units. For example, see [subtr\\_baseline\(\)](#). 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  $\log(1/T)$  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\\_baseline\(\)](#) for spectral background correction.

## Examples

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

*adj\_res*

*Normalization and conversion of spectral data*

## Description

`adj_res()` and `conform_res()` are helper functions to align wavenumbers in terms of their spectral resolution. `adj_neg()` converts numeric intensities  $y < 1$  into values  $\geq 1$ , keeping absolute differences between intensity values by shifting each value by the minimum intensity. `make_rel()` converts intensities  $y$  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_res(x, res = 1, fun = round)

conform_res(x, res = 5)

adj_neg(y, na.rm = FALSE)

mean_replace(y, na.rm = TRUE)

is_empty_vector(x)
```

**Arguments**

x	a numeric vector or an R object which is coercible to one by <code>as.vector(x, "numeric")</code> ; x should contain the spectral wavenumbers.
res	spectral resolution supplied to <code>fun</code> .
fun	the function to be applied to each element of <code>x</code> ; defaults to <code>round()</code> to round to a specific resolution <code>res</code> .
y	a numeric vector containing the spectral intensities.
na.rm	logical. Should missing values be removed?

**Details**

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

**Value**

`adj_res()` and `conform_res()` return a numeric vector with resolution-conformed wavenumbers. `adj_neg()` and `make_rel()` return numeric vectors with the normalized intensity data.

**Author(s)**

Win Cowger, Zacharias Steinmetz

**See Also**

`min()` and `round()`; `adj_intens()` for log transformation functions; `conform_spec()` for conforming wavenumbers of an OpenSpecy object to be matched with a reference library

**Examples**

```
adj_res(seq(500, 4000, 4), 5)
conform_res(seq(500, 4000, 4))
adj_neg(c(-1000, -1, 0, 1, 10))
```

---

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

---

**adj\_wave**

*Adjust wavelength to wavenumbers for Raman*

---

## Description

Functions for converting between wave\* units.

## Usage

```
adj_wave(x, ...)

## Default S3 method:
adj_wave(x, laser, ...)

## S3 method for class 'OpenSpecy'
adj_wave(x, laser, ...)
```

## Arguments

- x an OpenSpecy object with wavenumber units specified as wavelength in nm or a wavelength vector.
- laser the wavelength in nm of the Raman laser.
- ... additional arguments passed to submethods.

## Value

An OpenSpecy object with new units converted from wavelength to wavenumbers or a vector with the same conversion.

## Author(s)

Win Cowger, Zacharias Steinmetz

## Examples

```
data("raman_hdpe")
raman_wavelength <- raman_hdpe
raman_wavelength$wavenumber <- (-1*(raman_wavelength$wavenumber/10^7-1/530))^{(-1)}
adj_wave(raman_wavelength, laser = 530)
adj_wave(raman_wavelength$wavenumber, laser = 530)
```

---

as_OpenSpecy	<i>Create OpenSpecy objects</i>
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---

## Description

Functions to check if an object is an OpenSpecy, or coerce it if possible.

## Usage

```
as_OpenSpecy(x, ...)

## S3 method for class 'OpenSpecy'
as_OpenSpecy(x, session_id = FALSE, ...)

## S3 method for class 'list'
as_OpenSpecy(x, ...)

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

## S3 method for class 'data.frame'
as_OpenSpecy(x, colnames = list(wavenumber = NULL, spectra = NULL), ...)

## Default S3 method:
as_OpenSpecy(
  x,
  spectra,
  metadata = list(file_name = NULL, user_name = NULL, contact_info = NULL, organization =
    NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL, material_form =
    NULL, material_phase = NULL, material_producer = NULL, material_purity = NULL,
    material_quality = NULL, material_color = NULL, material_other = NULL, cas_number =
    NULL, instrument_used = NULL, instrument_accessories = NULL, instrument_mode = NULL,
    intensity_units = NULL, spectral_resolution = NULL, laser_light_used = NULL,
    number_of_accumulations = NULL,
    total_acquisition_time_s = NULL,
    data_processing_procedure = NULL, level_of_confidence_in_identification = NULL,
    other_info = NULL, license = "CC BY-NC"),
  attributes = list(intensity_unit = NULL, derivative_order = NULL, baseline = NULL,
    spectra_type = NULL),
  coords = "gen_grid",
  session_id = FALSE,
  comma_decimal = FALSE,
  ...
)
is_OpenSpecy(x)
```

```
check_OpenSpecy(x)
OpenSpecy(x, ...)
gen_grid(n)
```

### Arguments

x	depending on the method, a list with all OpenSpecy parameters, a vector with the wavenumbers for all spectra, or a data.frame with a full spectrum in the classic Open Specy format.
session_id	logical. Whether to add a session ID to the metadata. The session ID is based on current session info so metadata of the same spectra will not return equal if session info changes. Sometimes that is desirable.
colnames	names of the wavenumber column and spectra column, makes assumptions based on column names or placement if NULL.
spectra	spectral intensities formatted as a data.table with one column per spectrum.
metadata	metadata for each spectrum with one row per spectrum, see details.
attributes	a list of attributes describing critical aspects for interpreting the spectra. see details.
coords	spatial coordinates for the spectra.
comma_decimal	logical(1) whether commas may represent decimals.
n	number of spectra to generate the spatial coordinate grid with.
...	additional arguments passed to submethods.

### Details

as\_OpenSpecy() converts spectral datasets to a three part list; the first with a vector of the wavenumbers of the spectra, the second with a data.table of all spectral intensities ordered as columns, the third item is another data.table with any metadata the user provides or is harvested from the files themselves.

The metadata argument may contain a named list with the following details (\* = minimum recommended).

**file\_name\*** The file name, defaults to `basename()` if not specified  
**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., & Gerdts, G. (2018). Reference database design for the automated analysis of microplastic samples based on Fourier transform infrared (FTIR) spectroscopy. *Analytical and Bioanalytical Chemistry*. doi:10.1007/s00216-0181156x"  
**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, analytical standard, environmental sample  
**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  
**intensity\_units\*** Units of the intensity values for the spectrum, options transmittance, reflectance, absorbance  
**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/> for details). Any other creative commons license is allowed, for example, CC0 or CC BY  
**session\_id** A unique user and session identifier; populated automatically with paste(digest(Sys.info()), digest(sessionInfo()), sep = "/")  
**file\_id** A unique file identifier; populated automatically with digest(object[c("wavenumber", "spectra")])

The attributes argument may contain a named list with the following details, when set, they will be used to automate transformations and warning messages:

**intensity\_units** supported options include "absorbance", "transmittance", or "reflectance"  
**derivative\_order** supported options include "0", "1", or "2"  
**baseline** supported options include "raw" or "nobaseline"  
**spectra\_type** supported options include "ftir" or "raman"

## Value

`as_OpenSpecy()` and `OpenSpecy()` returns three part lists described in details. `is_OpenSpecy()` returns TRUE if the object is an OpenSpecy and FALSE if not. `gen_grid()` returns a `data.table` with x and y coordinates to use for generating a spatial grid for the spectra if one is not specified in the data.

**Author(s)**

Zacharias Steinmetz, Win Cowger

**See Also**

[read\\_spec\(\)](#) for reading OpenSpecy objects.

**Examples**

```
data("raman_hdpe")

# Inspect the spectra
raman_hdpe # see how OpenSpecy objects print.
raman_hdpe$wavenumber # look at just the wavenumbers of the spectra.
raman_hdpe$spectra # look at just the spectral intensities data.table.
raman_hdpe$metadata # look at just the metadata of the spectra.

# Creating a list and transforming to OpenSpecy
as_OpenSpecy(list(wavenumber = raman_hdpe$wavenumber,
                  spectra = raman_hdpe$spectra,
                  metadata = raman_hdpe$metadata[,-c("x", "y")]))

# If you try to produce an OpenSpecy using an OpenSpecy it will just return
# the same object.
as_OpenSpecy(raman_hdpe)

# Creating an OpenSpecy from a data.frame
as_OpenSpecy(x = data.frame(wavenumber = raman_hdpe$wavenumber,
                            spectra = raman_hdpe$spectra$intensity))

# Test that the spectrum is formatted as an OpenSpecy object.
is_OpenSpecy(raman_hdpe)
is_OpenSpecy(raman_hdpe$spectra)
```

[check\\_lib](#)

*Manage spectral libraries*

**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. Please see the references when using these libraries. These libraries are the accumulation of a massive amount of effort from independant groups and each should be attributed when you are using their data.

**Usage**

```

check_lib(
  type = c("derivative", "nobaseline", "raw", "medoid_derivative", "medoid_nobaseline",
          "model_derivative", "model_nobaseline"),
  path = "system",
  condition = "warning"
)

get_lib(
  type = c("derivative", "nobaseline", "raw", "medoid_derivative", "medoid_nobaseline",
          "model_derivative", "model_nobaseline"),
  path = "system",
  mode = "wb",
  revision = NULL,
  aws = FALSE,
  ...
)

load_lib(type, path = "system")

rm_lib(
  type = c("derivative", "nobaseline", "raw", "medoid_derivative", "medoid_nobaseline",
          "model_derivative", "model_nobaseline"),
  path = "system"
)

```

**Arguments**

type	library type to check/retrieve; defaults to c("derivative", "nobaseline", "raw", "medoid_derivative", "medoid_nobaseline", "model_derivative", "model_nobaseline") which reads everything.
path	where to save or look for local library files; defaults to "system" pointing to system.file("extdata", package = "OpenSpecy").
condition	determines if check_lib() should warn ("warning", the default) or throw and error ("error").
mode	see ?download.file for details on mode.
revision	revision number to use for libraries, revision numbers can be found at the osf repo ( <a href="https://osf.io/x7dpz/">https://osf.io/x7dpz/</a> ) by clicking the library then history, if NULL defaults to most recent. This allows exact version control.
aws	whether to source the files from AWS or OSF, default of FALSE is OSF.
...	further arguments passed to <a href="#">osf_download()</a> .

**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. `rm_lib()` removes the libraries from your computer.

### Value

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

### Author(s)

Zacharias Steinmetz, Win Cowger

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- Further contribution of spectra:** Suja Sukumaran (Thermo Fisher Scientific), Aline Carvalho, Jennifer Lynch (NIST), Claudia Cellia and Dora Mehn (JRC), Horiba Scientific, USDA Soil Characterization Data (<https://ncsslabdatamart.sc.egov.usda.gov>), Archaeometrielabor, and S.B. Engelsen (Royal Vet. and Agricultural University, Denmark). Kimmel Center data was collected and provided by Prof. Steven Weiner (Kimmel Center for Archaeological Science, Weizmann Institute of Science, Israel).

## Examples

```
## Not run:
check_lib("derivative")
get_lib("derivative")

spec_lib <- load_lib("derivative")

## End(Not run)
```

## Description

Functions for analyzing features, like particles, fragments, or fibers, in spectral map oriented OpenSpecy object.

## Usage

```
collapse_spec(x, ...)

## Default S3 method:
collapse_spec(x, ...)

## S3 method for class 'OpenSpecy'
collapse_spec(x, fun = median, column = "feature_id", ...)

def_features(x, ...)

## Default S3 method:
def_features(x, ...)

## S3 method for class 'OpenSpecy'
def_features(
  x,
  features,
  shape_kernel = c(3, 3),
  shape_type = "box",
  close = F,
  close_kernel = c(4, 4),
  close_type = "box",
  img = NULL,
  bottom_left = NULL,
  top_right = NULL,
  ...
)
```

## Arguments

<code>x</code>	an OpenSpecy object
<code>fun</code>	function name to collapse by.
<code>column</code>	column name in metadata to collapse by.
<code>features</code>	a logical vector or character vector describing which of the spectra are of features (TRUE) and which are not (FALSE). If a character vector is provided, it should represent the different feature types present in the spectra.
<code>shape_kernel</code>	the width and height of the area in pixels to search for connecting features, c(3,3) is typically used but larger numbers will smooth connections between particles more.
<code>shape_type</code>	character, options are for the shape used to find connections c("box", "disc", "diamond")

close	logical, whether a closing should be performed using the shape kernel before estimating components.
close_kernel	width and height of the area to close if using the close option.
close_type	character, options are for the shape used to find connections c("box", "disc", "diamond")
img	a file location where a visual image is that corresponds to the spectral image.
bottom_left	a two value vector specifying the x,y location in image pixels where the bottom left of the spectral map begins. y values are from the top down while x values are left to right.
top_right	a two value vector specifying the x,y location in the visual image pixels where the top right of the spectral map extent is. y values are from the top down while x values are left to right.
...	additional arguments passed to subfunctions.

## Details

`def_features()` accepts an OpenSpecy object and a logical or character vector describing which pixels correspond to particles. `collapse_spec()` takes an OpenSpecy object with particle-specific metadata (from `def_features()`) and collapses the spectra with a function intensities for each unique particle. It also updates the metadata with centroid coordinates, while preserving the feature information on area and Feret max.

## Value

An OpenSpecy object appended with metadata about the features or collapsed for the features. All units are in pixels. Metadata described below.

- x x coordinate of the pixel or centroid if collapsed
- y y coordinate of the pixel or centroid if collapsed
- feature\_id unique identifier of each feature
- area area in pixels of the feature
- perimeter perimeter of the convex hull of the feature
- feret\_min feret\_max divided by the area
- feret\_max largest dimension of the convex hull of the feature
- convex\_hull\_area area of the convex hull
- centroid\_x mean x coordinate of the feature
- centroid\_y mean y coordinate of the feature
- first\_x first x coordinate of the feature
- first\_y first y coordinate of the feature
- rand\_x random x coordinate from the feature
- rand\_y random y coordinate from the feature
- r if using visual imagery overlay, the red band value at that location
- g if using visual imagery overlay, the green band value at that location
- b if using visual imagery overlay, the blue band value at that location

**Author(s)**

Win Cowger, Zacharias Steinmetz

**Examples**

```
tiny_map <- read_extdata("CA_tiny_map.zip") |> read_any()
identified_map <- def_features(tiny_map, tiny_map$metadata$x == 0)
collapse_spec(identified_map)
```

conform\_spec

*Conform spectra to a standard wavenumber series*

**Description**

Spectra can be conformed to a standard suite of wavenumbers to be compared with a reference library or to be merged to other spectra.

**Usage**

```
conform_spec(x, ...)

## Default S3 method:
conform_spec(x, ...)

## S3 method for class 'OpenSpecy'
conform_spec(x, range = NULL, res = 5, allow_na = F, type = "interp", ...)
```

**Arguments**

x	a list object of class OpenSpecy.
range	a vector of new wavenumber values, can be just supplied as a min and max value.
res	spectral resolution adjusted to or NULL if the raw range should be used.
allow_na	logical; should NA values in places beyond the wavenumbers of the dataset be allowed?
type	the type of wavenumber adjustment to make. "interp" results in linear interpolation while "roll" conducts a nearest rolling join of the wavenumbers. "mean_up" only works when Spectra are being aggregated, we take the mean of the intensities within the wavenumber specified. This can maintain smaller peaks and make spectra more similar to it's less resolved relatives. mean_up option is still experimental.
...	further arguments passed to <a href="#">approx()</a>

**Value**

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

### Author(s)

Win Cowger, Zacharias Steinmetz

### See Also

[restrict\\_range\(\)](#) and [flatten\\_range\(\)](#) for adjusting wavenumber ranges; [subtr\\_baseline\(\)](#) for spectral background correction

### Examples

```
data("raman_hdpe")
conform_spec(raman_hdpe, c(1000, 2000))
```

---

cor_spec	<i>Identify and filter spectra</i>
----------	------------------------------------

---

### Description

`match_spec()` joins two OpenSpecy objects and their metadata based on similarity. `cor_spec()` correlates two OpenSpecy objects, typically one with knowns and one with unknowns. `ident_spec()` retrieves the top match values from a correlation matrix and formats them with metadata. `get_metadata()` retrieves metadata from OpenSpecy objects. `max_cor_named()` formats the top correlation values from a correlation matrix as a named vector. `filter_spec()` filters an Open Specy object. `fill_spec()` adds filler values to an OpenSpecy object where it doesn't have intensities. `os_similarity()` EXPERIMENTAL, returns a single similarity metric between two OpenSpecy objects based on the method used.

### Usage

```
cor_spec(x, ...)

## Default S3 method:
cor_spec(x, ...)

## S3 method for class 'OpenSpecy'
cor_spec(x, library, na.rm = T, conform = F, type = "roll", ...)

match_spec(x, ...)

## Default S3 method:
match_spec(x, ...)

## S3 method for class 'OpenSpecy'
match_spec(
  x,
  library,
```

```
na.rm = T,
conform = F,
type = "roll",
top_n = NULL,
order = NULL,
add_library_metadata = NULL,
add_object_metadata = NULL,
fill = NULL,
...
)

ident_spec(
  cor_matrix,
  x,
  library,
  top_n = NULL,
  add_library_metadata = NULL,
  add_object_metadata = NULL,
  ...
)
get_metadata(x, ...)

## Default S3 method:
get_metadata(x, ...)

## S3 method for class 'OpenSpecy'
get_metadata(x, logic, rm_empty = TRUE, ...)

max_cor_named(cor_matrix, na.rm = T)

filter_spec(x, ...)

## Default S3 method:
filter_spec(x, ...)

## S3 method for class 'OpenSpecy'
filter_spec(x, logic, ...)

ai_classify(x, ...)

## Default S3 method:
ai_classify(x, ...)

## S3 method for class 'OpenSpecy'
ai_classify(x, library, fill = NULL, ...)

fill_spec(x, ...)
```

```

## Default S3 method:
fill_spec(x, ...)

## S3 method for class 'OpenSpecy'
fill_spec(x, fill, ...)

os_similarity(x, ...)

## Default S3 method:
os_similarity(x, ...)

## S3 method for class 'OpenSpecy'
os_similarity(x, y, method = "hamming", na.rm = T, ...)

```

## Arguments

x	an OpenSpecy object, typically with unknowns.
library	an OpenSpecy or glmnet object representing the reference library of spectra or model to use in identification.
na.rm	logical; indicating whether missing values should be removed when calculating correlations. Default is TRUE.
conform	Whether to conform the spectra to the library wavenumbers or not.
type	the type of conformation to make returned by conform_spec()
top_n	integer; specifying the number of top matches to return. If NULL (default), all matches will be returned.
order	an OpenSpecy used for sorting, ideally the unprocessed one; NULL skips sorting.
add_library_metadata	name of a column in the library metadata to be joined; NULL if you don't want to join.
add_object_metadata	name of a column in the object metadata to be joined; NULL if you don't want to join.
fill	an OpenSpecy object with a single spectrum to be used to fill missing values for alignment with the AI classification.
cor_matrix	a correlation matrix for object and library, can be returned by cor_spec()
logic	a logical or numeric vector describing which spectra to keep.
rm_empty	logical; whether to remove empty columns in the metadata.
y	an OpenSpecy object to perform similarity search against x.
method	the type of similarity metric to return.
...	additional arguments passed cor().

**Value**

`match_spec()` and `ident_spec()` will return a [data.table-class](#)() containing correlations between spectra and the library. The table has three columns: `object_id`, `library_id`, and `match_val`. Each row represents a unique pairwise correlation between a spectrum in the object and a spectrum in the library. If `top_n` is specified, only the top `top_n` matches for each object spectrum will be returned. If `add_library_metadata` is `is.character`, the library metadata will be added to the output. If `add_object_metadata` is `is.character`, the object metadata will be added to the output. `filter_spec()` returns an OpenSpecy object. `fill_spec()` returns an OpenSpecy object. `cor_spec()` returns a correlation matrix. `get_metadata()` returns a [data.table-class](#)() with the metadata for columns which have information. `os_similarity()` returns a single numeric value representing the type of similarity metric requested. 'wavenumber' similarity is based on the proportion of wavenumber values that overlap between the two objects, 'metadata' is the proportion of metadata column names, 'hamming' is something similar to the hamming distance where we discretize all spectra in the OpenSpecy object by wavenumber intensity values and then relate the wavenumber intensity value distributions by mean difference in min-max normalized space. 'pca' tests the distance between the OpenSpecy objects in PCA space using the first 4 component values and calculating the max-range normalized distance between the mean components. The first two metrics are pretty straightforward and definitely ready to go, the 'hamming' and 'pca' metrics are pretty experimental but appear to be working under our current test cases.

**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**

```
data("test_lib")

unknown <- read_extdata("ftir_ldpe_soil.asp") |>
  read_any() |>
  conform_spec(range = test_lib$wavenumber,
               res = spec_res(test_lib)) |>
  process_spec()
  cor_spec(unknown, test_lib)

match_spec(unknown, test_lib, add_library_metadata = "sample_name",
           top_n = 1)
```

---

c_spec	<i>Manage spectral objects</i>
--------	--------------------------------

---

## Description

c\_spec() concatenates OpenSpecy objects. sample\_spec() samples spectra from an OpenSpecy object. merge\_map() merge two OpenSpecy objects from spectral maps.

## Usage

```
c_spec(x, ...)

## Default S3 method:
c_spec(x, ...)

## S3 method for class 'OpenSpecy'
c_spec(x, ...)

## S3 method for class 'list'
c_spec(x, range = NULL, res = 5, ...)

sample_spec(x, ...)

## Default S3 method:
sample_spec(x, ...)

## S3 method for class 'OpenSpecy'
sample_spec(x, size = 1, prob = NULL, ...)

merge_map(x, ...)

## Default S3 method:
merge_map(x, ...)

## S3 method for class 'OpenSpecy'
merge_map(x, ...)

## S3 method for class 'list'
merge_map(x, origins = NULL, ...)
```

## Arguments

- |       |  |
|-------|--|
| x     | a list of OpenSpecy objects or of file paths.  |
| range | a numeric providing your own wavenumber ranges or character argument called "common" to let c_spec() find the common wavenumber range of the supplied spectra. NULL will interpret the spectra having all the same wavenumber range. |

<code>res</code>	defaults to NULL, the resolution you want the output wavenumbers to be.
<code>size</code>	the number of spectra to sample.
<code>prob</code>	probabilities to use for the sampling.
<code>origins</code>	a list with 2 value vectors of x y coordinates for the offsets of each image.
<code>...</code>	further arguments passed to submethods.

**Value**

`c_spec()` and `sample_spec()` return `OpenSpecy` objects.

**Author(s)**

Zacharias Steinmetz, Win Cowger

**See Also**

[conform\\_spec\(\)](#) for conforming wavenumbers

**Examples**

```
# Concatenating spectra
spectra <- lapply(c(read_extdata("raman_hdpe.csv"),
                     read_extdata("ftir_ldpe_soil.asp")), read_any)
common <- c_spec(spectra, range = "common", res = 5)
range <- c_spec(spectra, range = c(1000, 2000), res = 5)

# Sampling spectra
tiny_map <- read_any(read_extdata("CA_tiny_map.zip"))
sampled <- sample_spec(tiny_map, size = 3)
```

**Description**

Methods to visualize and convert `OpenSpecy` objects.

**Usage**

```
## S3 method for class 'OpenSpecy'
head(x, ...)

## S3 method for class 'OpenSpecy'
print(x, ...)

## S3 method for class 'OpenSpecy'
```

```

plot(
  x,
  offset = 0,
  legend_var = NULL,
  pallet = rainbow,
  main = "Spectra Plot",
  xlab = "Wavenumber (1/cm)",
  ylab = "Intensity (a.u.)",
  ...
)

## S3 method for class 'OpenSpecy'
summary(object, ...)

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

## S3 method for class 'OpenSpecy'
as.data.table(x, ...)

```

## Arguments

x	an OpenSpecy object.
offset	Numeric value for vertical offset of each successive spectrum. Defaults to 1. If 0, all spectra share the same baseline.
legend_var	Character string naming a metadata column in x\$metadata that labels/colors each spectrum. If NULL, spectra won't be labeled.
pallet	The base R graphics color pallet function to use. If NULL will default to all black.
main	Plot text for title.
xlab	Plot x axis text
ylab	Plot y axis text
object	an OpenSpecy object.
...	further arguments passed to the respective default method.

## Details

head() shows the first few lines of an OpenSpecy object. print() prints the contents of an OpenSpecy object. plot() produces a [plot\(\)](#) of an OpenSpecy

## Value

head(), print(), and summary() return a textual representation of an OpenSpecy object. plot() and lines() return a plot. as.data.frame() and as.data.table() convert OpenSpecy objects into tabular data.

**Author(s)**

Zacharias Steinmetz, Win Cowger

**See Also**

[head\(\)](#), [print\(\)](#), [summary\(\)](#), [matplotlib\(\)](#), and [matlines\(\)](#), [as.data.frame\(\)](#), [as.data.table\(\)](#)

**Examples**

```
data("raman_hdpe")

# Printing the OpenSpecy object
print(raman_hdpe)

# Displaying the first few lines of the OpenSpecy object
head(raman_hdpe)

# Plotting the spectra
plot(raman_hdpe)
```

---

human\_ts

*Create human readable timestamps*

---

**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, Zacharias Steinmetz

**See Also**

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

## Examples

```
human_ts()
```

---

make\_rel

*Make spectral intensities relative*

---

## Description

`make_rel()` converts intensities  $x$  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

```
make_rel(x, ...)

## Default S3 method:
make_rel(x, na.rm = FALSE, ...)

## S3 method for class 'OpenSpecy'
make_rel(x, na.rm = FALSE, ...)
```

## Arguments

<code>x</code>	a numeric vector or an R OpenSpecy object
<code>na.rm</code>	logical. Should missing values be removed?
<code>...</code>	further arguments passed to <code>make_rel()</code> .

## Details

`make_rel()` is used to retain the relative height proportions between spectra while avoiding the large numbers that can result from some spectral instruments.

## Value

`make_rel()` return numeric vectors (if vector provided) or an OpenSpecy object with the normalized intensity data.

## Author(s)

Win Cowger, Zacharias Steinmetz

## See Also

`min()` and `round()`; `adj_intens()` for log transformation functions; `conform_spec()` for conforming wavenumbers of an OpenSpecy object to be matched with a reference library

## Examples

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

manage\_na

*Ignore or remove NA intensities*

## Description

Sometimes you want to keep or remove NA values in intensities to allow for spectra with varying shapes to be analyzed together or maintained in a single Open Specy object.

## Usage

```
manage_na(x, ...)

## Default S3 method:
manage_na(x, lead_tail_only = TRUE, ig = c(NA), ...)

## S3 method for class 'OpenSpecy'
manage_na(x, lead_tail_only = TRUE, ig = c(NA), fun, type = "ignore", ...)
```

## Arguments

x	a numeric vector or an R OpenSpecy object.
lead_tail_only	logical whether to only look at leading adn tailing values.
ig	character vector, values to ignore.
fun	the name of the function you want run, this is only used if the "ignore" type is chosen.
type	character of either "ignore" or "remove".
...	further arguments passed to fun.

## Value

manage\_na() return logical vectors of NA locations (if vector provided) or an OpenSpecy object with ignored or removed NA values.

## Author(s)

Win Cowger, Zacharias Steinmetz

## See Also

OpenSpecy object to be matched with a reference library `fill_spec()` can be used to fill NA values in Open Specy objects. `restrict_range()` can be used to restrict spectral ranges in other ways than removing NAs.

## Examples

```
manage_na(c(NA, -1, NA, 1, 10))
manage_na(c(NA, -1, NA, 1, 10), lead_tail_only = FALSE)
manage_na(c(NA, 0, NA, 1, 10), lead_tail_only = FALSE, ig = c(NA,0))
data(raman_hdpe)
raman_hdpe$spectra[[1]][1:10] <- NA

#would normally return all NA without na.rm = TRUE but doesn't here.
manage_na(raman_hdpe, fun = make_rel)

#will remove the first 10 values we set to NA
manage_na(raman_hdpe, type = "remove")
```

## Description

These functions generate heatmaps, spectral plots, and interactive plots for OpenSpecy data.

## Usage

```
plotly_spec(x, ...)

## Default S3 method:
plotly_spec(x, ...)

## S3 method for class 'OpenSpecy'
plotly_spec(
  x,
  x2 = NULL,
  line = list(color = "rgb(255, 255, 255)"),
  line2 = list(dash = "dot", color = "rgb(255,0,0)"),
  font = list(color = "#FFFFFF"),
  plot_bgcolor = "rgba(17, 0, 73, 0)",
  paper_bgcolor = "rgb(0, 0, 0)",
  showlegend = FALSE,
  make_rel = TRUE,
  ...
)

heatmap_spec(x, ...)

## Default S3 method:
heatmap_spec(x, ...)
```

```

## S3 method for class 'OpenSpecy'
heatmap_spec(
  x,
  z = NULL,
  sn = NULL,
  cor = NULL,
  min_sn = NULL,
  min_cor = NULL,
  select = NULL,
  font = list(color = "#FFFFFF"),
  plot_bgcolor = "rgba(17, 0, 73, 0)",
  paper_bgcolor = "rgb(0, 0, 0)",
  colorscale = "Viridis",
  showlegend = FALSE,
  type = "interactive",
  ...
)

interactive_plot(x, ...)

## Default S3 method:
interactive_plot(x, ...)

## S3 method for class 'OpenSpecy'
interactive_plot(
  x,
  x2 = NULL,
  select = NULL,
  line = list(color = "rgb(255, 255, 255)"),
  line2 = list(dash = "dot", color = "rgb(255,0,0)"),
  font = list(color = "#FFFFFF"),
  plot_bgcolor = "rgba(17, 0, 73, 0)",
  paper_bgcolor = "rgb(0, 0, 0)",
  colorscale = "Viridis",
  ...
)

```

## Arguments

- `x` an `OpenSpecy` object containing metadata and spectral data for the first group.
- `x2` an optional second `OpenSpecy` object containing metadata and spectral data for the second group.
- `line` list; `line` parameter for `x`; passed to [add\\_trace\(\)](#).
- `line2` list; `line` parameter for `x2`; passed to
- `font` list; passed to [layout\(\)](#).
- `plot_bgcolor` color value; passed to [layout\(\)](#).
- `paper_bgcolor` color value; passed to [layout\(\)](#).

showlegend	whether to show the legend passed to
make_rel	logical, whether to make the spectra relative or use the raw values
z	optional numeric vector specifying the intensity values for the heatmap. If not provided, the function will use the intensity values from the OpenSpecy object.
sn	optional numeric value specifying the signal-to-noise ratio threshold. If provided along with <code>min_sn</code> , regions with SNR below the threshold will be excluded from the heatmap.
cor	optional numeric value specifying the correlation threshold. If provided along with <code>min_cor</code> , regions with correlation below the threshold will be excluded from the heatmap.
min_sn	optional numeric value specifying the minimum signal-to-noise ratio for inclusion in the heatmap. Regions with SNR below this threshold will be excluded.
min_cor	optional numeric value specifying the minimum correlation for inclusion in the heatmap. Regions with correlation below this threshold will be excluded.
select	optional index of the selected spectrum to highlight on the heatmap.
colorscale	colorscale passed to <code>add_trace()</code> can be an array or one of "Blackbody", "Bluered", "Blues", "Cividis", "Earth", "Electric", "Greens", "Greys", "Hot", "Jet", "Picnic", "Portland", "Rainbow", "RdBu", "Reds", "Viridis", "YlGnBu", "YlOrRd".
type	specification for plot type either interactive or static <code>plot_ly()</code> .
...	further arguments passed to <code>plot_ly()</code> .

## Value

A `plotly` heatmap object displaying the OpenSpecy data. A subplot containing the heatmap and spectra plot. A `plotly` object displaying the spectra from the OpenSpecy object(s).

## Author(s)

Win Cowger, Zacharias Steinmetz

## Examples

```
data("raman_hdpe")
tiny_map <- read_extdata("CA_tiny_map.zip") |> read_zip()
plotly_spec(raman_hdpe)

heatmap_spec(tiny_map, z = tiny_map$metadata$y, showlegend = TRUE)

sample_spec(tiny_map, size = 12) |>
  interactive_plot(select = 2, x2 = raman_hdpe)
```

---

`process_spec`

*Process Spectra*

---

## Description

`process_spec()` is a monolithic wrapper function for all spectral processing steps.

## Usage

```
process_spec(x, ...)

## Default S3 method:
process_spec(x, ...)

## S3 method for class 'OpenSpecy'
process_spec(
  x,
  active = TRUE,
  adj_intens = FALSE,
  adj_intens_args = list(type = "none"),
  conform_spec = TRUE,
  conform_spec_args = list(range = NULL, res = 5, type = "interp"),
  restrict_range = FALSE,
  restrict_range_args = list(min = 0, max = 6000),
  flatten_range = FALSE,
  flatten_range_args = list(min = 2200, max = 2420),
  subtr_baseline = FALSE,
  subtr_baseline_args = list(type = "polynomial", degree = 8, raw = FALSE, baseline =
    NULL),
  smooth_intens = TRUE,
  smooth_intens_args = list(polynomial = 3, window = 11, derivative = 1, abs = TRUE),
  make_rel = TRUE,
  make_rel_args = list(na.rm = TRUE),
  ...
)
```

## Arguments

<code>x</code>	an OpenSpecy object.
<code>active</code>	logical; indicating whether to perform processing. If TRUE, the processing steps will be applied. If FALSE, the original data will be returned.
<code>adj_intens</code>	logical; describing whether to adjust the intensity units.
<code>adj_intens_args</code>	named list of arguments passed to <code>adj_intens()</code> .
<code>conform_spec</code>	logical; whether to conform the spectra to a new wavenumber range and resolution.

```

conform_spec_args
    named list of arguments passed to conform_spec().
restrict_range logical; indicating whether to restrict the wavenumber range of the spectra.
restrict_range_args
    named list of arguments passed to restrict_range().
flatten_range logical; indicating whether to flatten the range around the carbon dioxide region.
flatten_range_args
    named list of arguments passed to flatten_range().
subtr_baseline logical; indicating whether to subtract the baseline from the spectra.
subtr_baseline_args
    named list of arguments passed to subtr_baseline().
smooth_intens logical; indicating whether to apply a smoothing filter to the spectra.
smooth_intens_args
    named list of arguments passed to smooth_intens().
make_rel logical; if TRUE spectra are automatically normalized with make_rel().
make_rel_args named list of arguments passed to make_rel().
na.rm Whether to allow NA or set all NA values to
...
    further arguments passed to subfunctions.

```

## Value

`process_spec()` returns an OpenSpecy object with processed spectra based on the specified parameters.

## Examples

```

data("raman_hdpe")
plot(raman_hdpe)

# Process spectra with range restriction and baseline subtraction
process_spec(raman_hdpe,
             restrict_range = TRUE,
             restrict_range_args = list(min = 500, max = 3000),
             subtr_baseline = TRUE,
             subtr_baseline_args = list(type = "polynomial",
                                         polynomial = 8)) |>
  plot()

# Process spectra with smoothing and derivative
process_spec(raman_hdpe,
             smooth_intens = TRUE,
             smooth_intens_args = list(
               polynomial = 3,
               window = 11,
               derivative = 1
             )
           ) |>
  plot()

```

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

### Description

Raman spectrum of high-density polyethylene (HDPE) provided by Horiba Scientific.

### Format

An threepart list of class [OpenSpecy](#) containing:

wavenumber:	spectral wavenumbers [1/cm] (vector of 964 rows)
spectra:	absorbance values - (a <a href="#">data.table</a> with 964 rows and 1 column)
metadata:	spectral metadata

### Author(s)

Zacharias Steinmetz, Win Cowger

### References

Cowger W, Gray A, 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).

### Examples

```
data(raman_hdpe)
print(raman_hdpe)
```

read_any	<i>Read spectral data from multiple files</i>
----------	---

### Description

Wrapper functions for reading files in batch.

### Usage

```
read_any(file, c_spec = T, c_spec_args = list(range = NULL, res = NULL), ...)
read_many(file, ...)
read_zip(file, ...)
```

## Arguments

file	file to be read from or written to.
c_spec	logical, if multiple spectra should be concatenated or not. Multiple spectra will return a list if this is false.
c_spec_args	list of arguments passed to c_spec()
...	further arguments passed to the submethods.

## Details

`read_any()` provides a single function to quickly read in any of the supported formats, it assumes that the file extension will tell it how to process the spectra. `read_zip()` provides functionality for reading in spectral map files with ENVI file format or as individual files in a zip folder. If individual files, spectra are concatenated. `read_many()` provides functionality for reading multiple files in a character vector and will return a list.

## Value

All `read_*`() functions return OpenSpecy objects if a single spectrum or map is provided, otherwise they provide a list of Open Specys.

## Author(s)

Zacharias Steinmetz, Win Cowger

## See Also

[read\\_spec\(\)](#) for submethods. [c\\_spec\(\)](#) for combining lists of Open Specys.

## Examples

```
read_extdata("raman_hdpe.csv") |> read_any()
read_extdata("ftir_ldpe_soil.asp") |> read_any()
read_extdata("testdata_zipped.zip") |> read_many()
read_extdata("CA_tiny_map.zip") |> read_many()
```

---

read\_envi

*Read ENVI data*

---

## Description

This function allows ENVI data import.

**Usage**

```
read_envi(
  file,
  header = NULL,
  spectral_smooth = F,
  sigma = c(1, 1, 1),
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
    organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
    material_form = NULL, material_phase = NULL, material_producer = NULL,
    material_purity = NULL, material_quality = NULL, material_color = NULL,
    material_other = NULL, cas_number = NULL, instrument_used = NULL,
    instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
    laser_light_used = NULL, number_of_accumulations = NULL,
    total_acquisition_time_s = NULL, data_processing_procedure = NULL,
    level_of_confidence_in_identification = NULL, other_info = NULL, license =
    "CC BY-NC"),
  ...
)
```

**Arguments**

<code>file</code>	name of the binary file.
<code>header</code>	name of the ASCII header file. If <code>NULL</code> , the name of the header file is guessed by looking for a second file with the same basename as <code>file</code> but with <code>.hdr</code> extension.
<code>spectral_smooth</code>	logical value determines whether spectral smoothing will be performed.
<code>sigma</code>	if <code>spectral_smooth</code> then this option applies the 3d standard deviations for the <code>gaussianSmooth</code> function from the <code>mmand</code> package to describe how spectral smoothing occurs on each dimension. The first two dimensions are <code>x</code> and <code>y</code> , the third is the wavenumbers.
<code>metadata</code>	a named list of the metadata; see <a href="#">as_OpenSpecy()</a> for details.
<code>...</code>	further arguments passed to the submethods.

**Details**

ENVI data usually consists of two files, an ASCII header and a binary data file. The header contains all information necessary for correctly reading the binary file via [read.ENVI\(\)](#).

**Value**

An OpenSpecy object.

**Author(s)**

Zacharias Steinmetz, Claudia Beleites

## See Also

`read_spec()` for reading .y(a)ml, .json, or .rds (OpenSpecy) files; `read_text()`, `read_asp()`, `read_spa()`, `read_spc()`, and `read_jdx()` for text files, .asp, .spa, .spc, and .jdx formats, respectively; `read_opus()` for reading .0 (OPUS) files; `read_zip()` and `read_any()` for wrapper functions; `read.ENVI()` `gaussianSmooth()`

---

read\_opus

*Read spectral data from Bruker OPUS binary files*

---

## Description

Read file(s) acquired with a Bruker Vertex FTIR Instrument. This function is basically a fork of `opus_read()` from <https://github.com/pierreroudier/opusreader>.

## Usage

```
read_opus(  
  file,  
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,  
  organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,  
  material_form = NULL, material_phase = NULL, material_producer = NULL,  
  material_purity = NULL, material_quality = NULL, material_color = NULL,  
  material_other = NULL, cas_number = NULL, instrument_used = NULL,  
  instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,  
  laser_light_used = NULL, number_of_accumulations = NULL,  
  
  total_acquisition_time_s = NULL, data_processing_procedure = NULL,  
  level_of_confidence_in_identification = NULL, other_info = NULL, license =  
  "CC BY-NC"),  
  type = "spec",  
  digits = 1L,  
  atm_comp_minus4offset = FALSE  
)
```

## Arguments

file	character vector with path to file(s).
metadata	a named list of the metadata; see <code>as_OpenSpecy()</code> for details.
type	character vector of spectra types to extract from OPUS binary file. Default is "spec", which will extract the final spectra, e.g. expressed in absorbance (named AB in Bruker OPUS programs). Possible additional values for the character vector supplied to type are "spec_no_atm_comp" (spectrum of the sample without compensation for atmospheric gases, water vapor and/or carbon dioxide), "sc_sample" (single channel spectrum of the sample measurement), "sc_ref" (single channel spectrum of the reference measurement), "ig_sample" (interferogram of the sample measurement) and "ig_ref" (interferogram of the reference measurement).

<code>digits</code>	Integer that specifies the number of decimal places used to round the wavenumbers (values of x-variables).
<code>atm_comp_minus4offset</code>	Logical whether spectra after atmospheric compensation are read with an offset of -4 bytes from Bruker OPUS files; default is FALSE.

## Details

The type of spectra returned by the function when using `type = "spec"` depends on the setting of the Bruker instrument: typically, it can be either absorbance or reflectance.

The type of spectra to extract from the file can also use Bruker's OPUS software naming conventions, as follows:

- ScSm corresponds to `sc_sample`
- ScRf corresponds to `sc_ref`
- IgSm corresponds to `ig_sample`
- IgRf corresponds to `ig_ref`

## Value

An OpenSpecy object.

## Author(s)

Philipp Baumann, Zacharias Steinmetz, Win Cowger

## See Also

[read\\_spec\(\)](#) for reading .y(a)m1, .json, or .rds (OpenSpecy) files; [read\\_text\(\)](#), [read\\_asp\(\)](#), [read\\_spa\(\)](#), [read\\_spc\(\)](#), and [read\\_jdx\(\)](#) for text files, .asp, .spa, .spa, .spc, and .jdx formats, respectively; [read\\_text\(\)](#) for reading .dat (ENVI) files; [read\\_zip\(\)](#) and [read\\_any\(\)](#) for wrapper functions; [read\\_opus\\_raw\(\)](#);

## Examples

```
read_extdata("ftir_ps.0") |> read_opus()
```

`read_opus_raw`

*Read a Bruker OPUS spectrum binary raw string*

## Description

Read single binary acquired with an Bruker Vertex FTIR Instrument

## Usage

```
read_opus_raw(rw, type = "spec", atm_comp_minus4offset = FALSE)
```

## Arguments

<code>rw</code>	a raw vector
<code>type</code>	character vector of spectra types to extract from OPUS binary file. Default is "spec", which will extract the final spectra, e.g. expressed in absorbance (named AB in Bruker OPUS programs). Possible additional values for the character vector supplied to type are "spec_no_atm_comp" (spectrum of the sample without compensation for atmospheric gases, water vapor and/or carbon dioxide), "sc_sample" (single channel spectrum of the sample measurement), "sc_ref" (single channel spectrum of the reference measurement), "ig_sample" (interferogram of the sample measurement) and "ig_ref" (interferogram of the reference measurement).
<code>atm_comp_minus4offset</code>	logical; whether spectra after atmospheric compensation are read with an offset of -4 bytes from Bruker OPUS files. Default is FALSE.

## Details

The type of spectra returned by the function when using `type = "spec"` depends on the setting of the Bruker instrument: typically, it can be either absorbance or reflectance.

The type of spectra to extract from the file can also use Bruker's OPUS software naming conventions, as follows:

- ScSm corresponds to `sc_sample`
- ScRf corresponds to `sc_ref`
- IgSm corresponds to `ig_sample`
- IgRf corresponds to `ig_ref`

## Value

A list of 10 elements:

`metadata` a `data.frame` containing metadata from the OPUS file.

`spec` if "spec" was requested in the `type` option, a matrix of the spectrum of the sample (otherwise set to NULL).

`spec_no_atm_comp` if "spec\_no\_atm\_comp" was requested in the `type` option, a matrix of the spectrum of the sample without atmospheric compensation (otherwise set to NULL).

`sc_sample` if "sc\_sample" was requested in the `type` option, a matrix of the single channel spectrum of the sample (otherwise set to NULL).

`sc_ref` if "sc\_ref" was requested in the `type` option, a matrix of the single channel spectrum of the reference (otherwise set to NULL).

`ig_sample` if "ig\_sample" was requested in the `type` option, a matrix of the interferogram of the sample (otherwise set to NULL).

`ig_ref` if "ig\_ref" was requested in the `type` option, a matrix of the interferogram of the reference (otherwise set to NULL).

wavenumbers if "spec" or "spec\_no\_atm\_comp" was requested in the type option, a numeric vector of the wavenumbers of the spectrum of the sample (otherwise set to NULL).

wavenumbers\_sc\_sample if "sc\_sample" was requested in the type option, a numeric vector of the wavenumbers of the single channel spectrum of the sample (otherwise set to NULL).

wavenumbers\_sc\_ref if "sc\_ref" was requested in the type option, a numeric vector of the wavenumbers of the single channel spectrum of the reference (otherwise set to NULL).

### Author(s)

Philipp Baumann and Pierre Roudier

### See Also

[read\\_opus\(\)](#)

*read\_text*

*Read spectral data*

### Description

Functions for reading spectral data from external file types. Currently supported reading formats are .csv and other text files, .asp, .spa, .spc, .xyz, and .jdx. Additionally, .0 (OPUS) and .dat (ENVI) files are supported via [read\\_opus\(\)](#) and [read\\_envi\(\)](#), respectively. [read\\_zip\(\)](#) takes any of the files listed above. Note that proprietary file formats like .0, .asp, and .spa are poorly supported but will likely still work in most cases.

### Usage

```
read_text(
  file,
  colnames = NULL,
  method = "fread",
  comma_decimal = TRUE,
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
    organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
    material_form = NULL, material_phase = NULL, material_producer = NULL,
    material_purity = NULL, material_quality = NULL, material_color = NULL,
    material_other = NULL, cas_number = NULL, instrument_used = NULL,
    instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
    laser_light_used = NULL, number_of_accumulations = NULL,
    total_acquisition_time_s = NULL, data_processing_procedure = NULL,
    level_of_confidence_in_identification = NULL, other_info = NULL, license =
    "CC BY-NC"),
  ...
)
```

```
read_asp(
  file,
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
  organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
  material_form = NULL, material_phase = NULL, material_producer = NULL,
  material_purity = NULL, material_quality = NULL, material_color = NULL,
  material_other = NULL, cas_number = NULL, instrument_used = NULL,
  instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
  laser_light_used = NULL, number_of_accumulations = NULL,
  total_acquisition_time_s = NULL, data_processing_procedure = NULL,
  level_of_confidence_in_identification = NULL, other_info = NULL, license =
  "CC BY-NC"),
  ...
)

read_spa(
  file,
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
  organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
  material_form = NULL, material_phase = NULL, material_producer = NULL,
  material_purity = NULL, material_quality = NULL, material_color = NULL,
  material_other = NULL, cas_number = NULL, instrument_used = NULL,
  instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
  laser_light_used = NULL, number_of_accumulations = NULL,
  total_acquisition_time_s = NULL, data_processing_procedure = NULL,
  level_of_confidence_in_identification = NULL, other_info = NULL, license =
  "CC BY-NC"),
  ...
)

read_spc(
  file,
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
  organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
  material_form = NULL, material_phase = NULL, material_producer = NULL,
  material_purity = NULL, material_quality = NULL, material_color = NULL,
  material_other = NULL, cas_number = NULL, instrument_used = NULL,
  instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
  laser_light_used = NULL, number_of_accumulations = NULL,
  total_acquisition_time_s = NULL, data_processing_procedure = NULL,
  level_of_confidence_in_identification = NULL, other_info = NULL, license =
  "CC BY-NC"),
  ...
)
```

```

read_jdx(
  file,
  metadata = list(file_name = basename(file), user_name = NULL, contact_info = NULL,
    organization = NULL, citation = NULL, spectrum_type = NULL, spectrum_identity = NULL,
    material_form = NULL, material_phase = NULL, material_producer = NULL,
    material_purity = NULL, material_quality = NULL, material_color = NULL,
    material_other = NULL, cas_number = NULL, instrument_used = NULL,
    instrument_accessories = NULL, instrument_mode = NULL, spectral_resolution = NULL,
    laser_light_used = NULL, number_of_accumulations = NULL,
    total_acquisition_time_s = NULL, data_processing_procedure = NULL,
    level_of_confidence_in_identification = NULL, other_info = NULL, license =
    "CC BY-NC"),
  ...
)

read_extdata(file = NULL)

```

## Arguments

file	file to be read from or written to.
colnames	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 <code>fread()</code> but <code>read.csv()</code> works as well.
comma_decimal	logical(1) whether commas may represent decimals.
metadata	a named list of the metadata; see <code>as_OpenSpecy()</code> for details.
...	further arguments passed to the submethods.

## Details

`read_spc()` and `read_jdx()` are wrappers around the functions provided by the `hyperSpec`. Other functions have been adapted various online sources. Metadata is harvested if possible. 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\\_spec\(\)](#) for reading .y(a)ml, .json, or .rds (OpenSpecy) files; [read\\_opus\(\)](#) for reading .0 (OPUS) files; [read\\_envi\(\)](#) for reading .dat (ENVI) files; [read\\_zip\(\)](#) and [read\\_any\(\)](#) for wrapper functions; [read.jdx\(\)](#); [read.spc\(\)](#)

**Examples**

```
read_extdata("raman_hdpe.csv") |> read_text()
read_extdata("raman_atacamit.spc") |> read_spc()
read_extdata("ftir_ldpe_soil.asp") |> read_asp()
read_extdata("testdata_zipped.zip") |> read_zip()
```

---

**restrict\_range***Range restriction and flattening for spectra*

---

**Description**

`restrict_range()` restricts wavenumber ranges to user specified values. Multiple ranges can be specified by inputting a series of max and min values in order. `flatten_range()` will flatten ranges of the spectra that should have no peaks. Multiple ranges can be specified by inputting the series of max and min values in order.

**Usage**

```
restrict_range(x, ...)

## Default S3 method:
restrict_range(x, ...)

## S3 method for class 'OpenSpecy'
restrict_range(x, min, max, make_rel = TRUE, ...)

flatten_range(x, ...)

## Default S3 method:
flatten_range(x, ...)

## S3 method for class 'OpenSpecy'
flatten_range(x, min = 2200, max = 2400, make_rel = TRUE, ...)
```

**Arguments**

- |     |   |
|-----|---|
| x   | an OpenSpecy object.                                      |
| min | a vector of minimum values for the range to be flattened. |
| max | a vector of maximum values for the range to be flattened. |

<code>make_rel</code>	logical; should the output intensities be normalized to the range [0, 1] using <code>make_rel()</code> function?
...	additional arguments passed to subfunctions; currently not in use.

**Value**

An OpenSpecy object with the spectral intensities within specified ranges restricted or flattened.

**Author(s)**

Win Cowger, Zacharias Steinmetz

**See Also**

[conform\\_spec\(\)](#) for conforming wavenumbers to be matched with a reference library; [adj\\_intens\(\)](#) for log transformation functions; [min\(\)](#) and [round\(\)](#)

**Examples**

```
test_noise <- as_OpenSpecy(x = seq(400,4000, by = 10),
                           spectra = data.frame(intensity = rnorm(361)))
plot(test_noise)

restrict_range(test_noise, min = 1000, max = 2000)

flattened_intensities <- flatten_range(test_noise, min = c(1000, 2000),
                                         max = c(1500, 2500))
plot(flattened_intensities)
```

**run\_app**

*Run Open Specy app*

**Description**

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

**Usage**

```
run_app(path = "system", log = TRUE, ref = "main", test_mode = FALSE, ...)
```

**Arguments**

<code>path</code>	to store the downloaded app files; defaults to "system" pointing to <code>system.file(package = "OpenSpecy")</code> .
<code>log</code>	logical; enables/disables logging to <code>tempdir()</code>
<code>ref</code>	git reference; could be a commit, tag, or branch name. Defaults to "main". Only change this in case of errors.
<code>test_mode</code>	logical; for internal testing only.
...	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 [runGitHub\(\)](#).

**Author(s)**

Zacharias Steinmetz

**See Also**

[runGitHub\(\)](#)

**Examples**

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

---

sig\_noise

*Calculate signal and noise metrics for OpenSpecy objects*

---

**Description**

This function calculates common signal and noise metrics for OpenSpecy objects.

**Usage**

```
sig_noise(x, ...)  
  
## Default S3 method:  
sig_noise(x, ...)  
  
## S3 method for class 'OpenSpecy'  
sig_noise(  
  x,  
  metric = "run_sig_over_noise",  
  na.rm = TRUE,  
  prob = 0.5,  
  step = 20,  
  breaks = seq(min(unlist(x$spectra)), max(unlist(x$spectra)), length =  
    ((nrow(x$spectra)^(1/3)) * (max(unlist(x$spectra)) - min(unlist(x$spectra))))/(2 *  
    IQR(unlist(x$spectra)))),
```

```

sig_min = NULL,
sig_max = NULL,
noise_min = NULL,
noise_max = NULL,
abs = T,
spatial_smooth = F,
sigma = c(1, 1),
threshold = NULL,
...
)

```

## Arguments

<b>x</b>	an OpenSpecy object.
<b>metric</b>	character; specifying the desired metric to calculate. Options include "sig" (mean intensity), "noise" (standard deviation of intensity), "sig_times_noise" (absolute value of signal times noise), "sig_over_noise" (absolute value of signal / noise), "run_sig_over_noise" (absolute value of signal / noise where signal is estimated as the max intensity and noise is estimated as the height of a low intensity region.), "log_tot_sig" (sum of the inverse log intensities, useful for spectra in log units), "tot_sig" (sum of intensities), or "entropy" (Shannon entropy of intensities)..
<b>na.rm</b>	logical; indicating whether missing values should be removed when calculating signal and noise. Default is TRUE.
<b>prob</b>	numeric single value; the probability to retrieve for the quantile where the noise will be interpreted with the run_sig_over_noise option.
<b>step</b>	numeric; the step size of the region to look for the run_sig_over_noise option.
<b>breaks</b>	numeric; the number or positions of the breaks for entropy calculation. Defaults to infer a decent value from the data.
<b>sig_min</b>	numeric; the minimum wavenumber value for the signal region.
<b>sig_max</b>	numeric; the maximum wavenumber value for the signal region.
<b>noise_min</b>	numeric; the minimum wavenumber value for the noise region.
<b>noise_max</b>	numeric; the maximum wavenumber value for the noise region.
<b>abs</b>	logical; whether to return the absolute value of the result
<b>spatial_smooth</b>	logical; whether to spatially smooth the sig/noise using the xy coordinates and a gaussian smoother.
<b>sigma</b>	numeric; two value vector describing standard deviation for smoother in each dimension, y is specified first followed by x, should be the same for each in most cases.
<b>threshold</b>	numeric; if NULL, no threshold is set, otherwise use a numeric value to set the target threshold which true signal or noise should be above. The function will return a logical value instead of numeric if a threshold is set.
<b>...</b>	further arguments passed to subfunctions; currently not used.

**Value**

A numeric vector containing the calculated metric for each spectrum in the OpenSpecy object or logical value if threshold is set describing if the numbers where above or equal to (TRUE) the threshold.

**See Also**

[restrict\\_range\(\)](#)

**Examples**

```
data("raman_hdpe")

sig_noise(raman_hdpe, metric = "sig")
sig_noise(raman_hdpe, metric = "noise")
sig_noise(raman_hdpe, metric = "sig_times_noise")
```

---

smooth\_intens

*Smooth spectral intensities*

---

**Description**

This smoother can enhance the signal to noise ratio of the data using a Savitzky-Golay or Whittaker-Henderson filter.

**Usage**

```
smooth_intens(x, ...)

## Default S3 method:
smooth_intens(x, ...)

## S3 method for class 'OpenSpecy'
smooth_intens(
  x,
  polynomial = 3,
  window = 11,
  derivative = 1,
  abs = TRUE,
  lambda = 1600,
  d = 2,
  type = "sg",
  lag = 2,
  make_rel = TRUE,
  ...
)
```

```

calc_window_points(x, ...)

## Default S3 method:
calc_window_points(x, wavenum_width = 70, ...)

## S3 method for class 'OpenSpecy'
calc_window_points(x, wavenum_width = 70, ...)

```

## Arguments

<code>x</code>	an object of class OpenSpecy or vector for <code>calc_window_points()</code> .
<code>polynomial</code>	polynomial order for the filter
<code>window</code>	number of data points in the window, filter length (must be odd).
<code>derivative</code>	the derivative order if you want to calculate the derivative. Zero (default) is no derivative.
<code>abs</code>	logical; whether you want to calculate the absolute value of the resulting output.
<code>lambda</code>	smoothing parameter for Whittaker-Henderson smoothing, 50 results in rough smoothing and $10^4$ results in a high level of smoothing.
<code>d</code>	order of differences to use for Whittaker-Henderson smoothing, typically set to 2.
<code>type</code>	the type of smoothing to use "wh" for Whittaker-Henerson or "sg" for Savitzky-Golay.
<code>lag</code>	the lag to use for the numeric derivative calculation if using Whittaker-Henderson. Greater values lead to smoother derivatives, 1 or 2 is common.
<code>make_rel</code>	logical; if TRUE spectra are automatically normalized with <code>make_rel()</code> .
<code>wavenum_width</code>	the width of the window you want in wavenumbers.
<code>...</code>	further arguments passed to <code>sgolay()</code> .

## Details

For Savitzky-Golay 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. For Whittaker-Henderson, the code is largely based off of the `whittaker()` function in the pracma package. In general Whittaker-Henderson is expected to be slower but more robust than Savitzky-Golay.

## Value

`smooth_intens()` returns an OpenSpecy object.

`calc_window_points()` returns a single numeric vector object of the number of points needed to fill the window and can be passed to `smooth_intens()`. For many applications, this is more reusable than specifying a static number of points.

## 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)  
  
smooth_intens(raman_hdpe, window = calc_window_points(x = raman_hdpe, wavenum_width = 70))  
  
smooth_intens(raman_hdpe, lambda = 1600, d = 2, lag = 2, type = "wh")
```

---

spatial\_smooth      *Spatial Smoothing of OpenSpecy Objects*

---

## Description

Applies spatial smoothing to an OpenSpecy object using a Gaussian filter.

## Usage

```
spatial_smooth(x, sigma = c(1, 1, 1), ...)
```

## Arguments

- |       |  |
|-------|--|
| x     | an OpenSpecy object.   |
| sigma | a numeric vector specifying the standard deviations for the Gaussian kernel in the x and y dimensions, respectively. |
| ...   | further arguments passed to or from other methods.   |

## Details

This function performs spatial smoothing on the spectral data in an OpenSpecy object. It assumes that the spatial coordinates are provided in the `metadata` element of the object, specifically in the `x` and `y` columns, and that there is a `col_id` column in `metadata` that matches the column names in the `spectra` data.table.

## Value

An OpenSpecy object with smoothed spectra.

**Author(s)**

Win Cowger

**See Also**

[as\\_OpenSpecy\(\)](#), [gaussianSmooth\(\)](#)

*spec\_res*

*Spectral resolution*

**Description**

Helper function for calculating the spectral resolution from wavenumber data.

**Usage**

```
spec_res(x, ...)

## Default S3 method:
spec_res(x, ...)

## S3 method for class 'OpenSpecy'
spec_res(x, ...)
```

**Arguments**

x	a numeric vector with wavenumber data or an OpenSpecy object.
...	further arguments passed to subfunctions; currently not used.

**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)
```

---

split_spec	<i>Split Open Specy objects</i>
------------	---------------------------------

---

## Description

Convert a list of Open Specy objects with any number of spectra into a list of Open Specy objects with one spectrum each.

## Usage

```
split_spec(x)
```

## Arguments

x	a list of OpenSpecy objects
---	-----------------------------

## Details

Function will accept a list of Open Specy objects of any length and will split them to their individual components. For example a list of two objects, an Open Specy with only one spectrum and an Open Specy with 50 spectra will return a list of length 51 each with Open Specy objects that only have one spectrum.

## Value

A list of Open Specy objects each with 1 spectrum.

## Author(s)

Zacharias Steinmetz, Win Cowger

## See Also

[c\\_spec\(\)](#) for combining OpenSpecy objects. [collapse\\_spec\(\)](#) for summarizing OpenSpecy objects.

## Examples

```
data("test_lib")
data("raman_hdpe")
listed <- list(test_lib, raman_hdpe)
test <- split_spec(listed)
test2 <- split_spec(list(test_lib))
```

---

**subtr\_baseline***Automated background subtraction for spectral data*

---

**Description**

This baseline correction routine iteratively finds the baseline of a spectrum using polynomial fitting methods or accepts a manual baseline.

**Usage**

```
subtr_baseline(x, ...)

## Default S3 method:
subtr_baseline(
  x,
  y,
  type = "polynomial",
  degree = 8,
  raw = FALSE,
  full = T,
  remove_peaks = T,
  refit_at_end = F,
  crop_boundaries = F,
  iterations = 10,
  peak_width_mult = 3,
  termination_diff = 0.05,
  degree_part = 2,
  bl_x = NULL,
  bl_y = NULL,
  make_rel = TRUE,
  ...
)

## S3 method for class 'OpenSpecy'
subtr_baseline(
  x,
  type = "polynomial",
  degree = 8,
  raw = FALSE,
  full = T,
  remove_peaks = T,
  refit_at_end = F,
  crop_boundaries = F,
  iterations = 10,
  peak_width_mult = 3,
  termination_diff = 0.05,
  degree_part = 2,
```

```
baseline = list(wavenumber = NULL, spectra = NULL),
make_rel = TRUE,
...
)
```

## Arguments

x	a list object of class OpenSpecy or a vector of wavenumbers.
y	a vector of spectral intensities.
type	one of "polynomial" or "manual" depending on the desired baseline correction method.
degree	the degree of the full spectrum polynomial. Must be less than the number of unique points when raw is FALSE. Typically, a good fit can be found with an 8th order polynomial.
raw	if TRUE, use raw and not orthogonal polynomials.
full	logical, whether to use the full spectrum as in "imodpoly" or to partition as in "smodpoly".
remove_peaks	logical, whether to remove peak regions during first iteration.
refit_at_end	logical, whether to refit a polynomial to the end result (TRUE) or to use linear approximation.
crop_boundaries	logical, whether to smartly crop the boundaries to match the spectra based on peak proximity.
iterations	the number of iterations for automated baseline correction.
peak_width_mult	scaling factor for the width of peak detection regions.
termination_diff	scaling factor for the ratio of difference in residual standard deviation to terminate iterative fitting with.
degree_part	the degree of the polynomial for "smodpoly". Must be less than the number of unique points.
bl_x	a vector of wavenumbers for the baseline.
bl_y	a vector of spectral intensities for the baseline.
make_rel	logical; if TRUE, spectra are automatically normalized with <code>make_rel()</code> .
baseline	an OpenSpecy object containing the baseline data to be subtracted (only for "manual").
...	further arguments passed to <code>poly()</code> or "smodpoly" parameters.

## Details

This function supports two types of "polynomial" automated baseline correction with options. Default settings are closest to "imodpoly" for iterative polynomial fitting based on Zhao et al. (2007). Additionally options recommended by "smodpoly" for segmented iterative polynomial fitting with enhanced peak detection from the S-Modpoly algorithm (<https://github.com/jackma123-rgb/S-Modpoly>), and "manual" for applying a user-provided baseline.

**Value**

`subtr_baseline()` 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.

Jackma123 (2023). S-Modpoly: Segmented modified polynomial fitting for spectral baseline correction. *GitHub Repository*. Retrieved from <https://github.com/jackma123-rgb/S-Modpoly>.

**See Also**

`poly()`; `smooth_intens()`

**Examples**

```
data("raman_hdpe")

# Use polynomial
subtr_baseline(raman_hdpe, type = "polynomial", degree = 8)

subtr_baseline(raman_hdpe, type = "polynomial", iterations = 5)

# Use manual
bl <- raman_hdpe
bl$spectra$intensity <- bl$spectra$intensity / 2
subtr_baseline(raman_hdpe, type = "manual", baseline = bl)
```

---

**Description**

Reference library with 29 FTIR and 28 Raman spectra used for examples and internal testing.

**Format**

An OpenSpecy object; `sample_name` is the class of the spectra.

**Author(s)**

Win Cowger

**Examples**

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

---

write_spec	<i>Read and write spectral data</i>
------------	-------------------------------------

---

**Description**

Functions for reading and writing spectral data to and from OpenSpecy format. OpenSpecy objects are lists with components wavenumber, spectra, and metadata. Currently supported formats are .y(a)ml, .json, .csv, or .rds.

**Usage**

```
write_spec(x, ...)

## Default S3 method:
write_spec(x, ...)

## S3 method for class 'OpenSpecy'
write_spec(x, file, method = NULL, digits =getOption("digits"), ...)

read_spec(file, method = NULL, ...)

as_hyperSpec(x)
```

**Arguments**

- |        |   |
|--------|---|
| x      | an object of class <a href="#">OpenSpecy</a> .  |
| file   | file path to be read from or written to.  |
| method | optional; function to be used as a custom reader or writer. Defaults to the appropriate function based on the file extension. |
| digits | number of significant digits to use when formatting numeric values; defaults to <a href="#">getOption("digits")</a> .         |
| ...    | further arguments passed to the submethods.   |

**Details**

Due to floating point number errors there may be some differences in the precision of the numbers returned if using multiple devices for .json and .yaml files but the numbers should be nearly identical. [readRDS\(\)](#) should return the exact same object every time.

**Value**

`read_spec()` reads data formatted as an `OpenSpecy` object and returns a list object of class `OpenSpecy` containing spectral data. `write_spec()` writes a file for an object of class `OpenSpecy` containing spectral data. `as_hyperspec()` converts an `OpenSpecy` object to a `hyperSpec-class` object.

**Author(s)**

Zacharias Steinmetz, Win Cowger

**See Also**

`OpenSpecy()`; `read_text()`, `read_asp()`, `read_spa()`, `read_spc()`, and `read_jdx()` for text files, .asp, .spa, .spa, .spc, and .jdx formats, respectively; `read_zip()` and `read_any()` for wrapper functions; `saveRDS()`; `readRDS()`; `write_yaml()`; `read_yaml()`; `write_json()`; `read_json()`;

**Examples**

```
read_extdata("raman_hdpe.yml") |> read_spec()
read_extdata("raman_hdpe.json") |> read_spec()
read_extdata("raman_hdpe.rds") |> read_spec()
read_extdata("raman_hdpe.csv") |> read_spec()

## Not run:
data(raman_hdpe)
write_spec(raman_hdpe, "raman_hdpe.yml")
write_spec(raman_hdpe, "raman_hdpe.json")
write_spec(raman_hdpe, "raman_hdpe.rds")
write_spec(raman_hdpe, "raman_hdpe.csv")

# Convert an OpenSpecy object to a hyperSpec object
hyper <- as_hyperSpec(raman_hdpe)

## End(Not run)
```

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