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LumiSpy is a Python package extending the functionality for multi-dimensional data analysis provided by the HyperSpy library. It is aimed at helping with the analysis of luminescence spectroscopy data (cathodoluminescence, photoluminescence, electroluminescence, Raman, SNOM).

Check out the Installation section for further information, including how to start using this project.

Complementing this documentation, the LumiSpy Demos repository contains curated Jupyter notebooks to provide tutorials and exemplary workflows.

**Note:** This project is under active development. Everyone is welcome to contribute. Please read our (see Contributing) guidelines and get started!
1.1 Installation

To install LumiSpy, you have the following options (independent of the operating system you use):

1. LumiSpy is included in the HyperSpy Bundle, a standalone program that includes a python distribution and all relevant libraries (recommended if you do not use python for anything else).

2. Installation using conda (recommended if you are also working with other python packages).

3. Installation using pip.

4. Installing the development version from GitHub. Refer to the appropriate section in the HyperSpy user guide (replacing hyperspy by lumispy).

1.1.1 Installation using conda

Follow these 3 steps to install LumiSpy using conda and start using it.

1. Creating a conda environment

LumiSpy requires Python 3 and conda – we suggest using the Python 3 version of Miniconda.

We recommend creating a new environment for the LumiSpy package (or installing it in the HyperSpy environment, if you have one already). To create a new environment:

1. Load the anaconda prompt.

2. Run the following command:

   (base) conda create -n lumispy -y

2. Installing the package in the new environment

Now activate the LumiSpy environment and install the package from conda-forge:

   (base) conda activate lumispy
   (lumispy) conda install -c conda-forge lumispy -y

Required dependencies will be installed automatically.

Installation is completed! To start using it, check the next section.
3. Getting Started

To get started using LumiSpy, especially if you are unfamiliar with Python, we recommend using Jupyter notebooks. Having installed LumiSpy as above, a Jupyter notebook can be installed and opened using the following commands entered into an anaconda prompt (from scratch):

(base) conda activate lumispy
(lumispy) conda install -c conda-forge jupyterlab -y
(lumispy) jupyter lab

Tutorials and exemplary workflows have been curated as a series of Jupyter notebooks that you can work through and modify to perform many common analyses.

1.1.2 Installation using pip

Alternatively, you can also find LumiSpy in the Python Package Index (PyPI) and install it using (requires pip):

```
pip install lumispy
```

Required dependencies will be installed automatically.

1.1.3 Updating the package

Using conda:

```
conda update lumispy -c conda-forge
```

Using pip:

```
pip install lumispy --upgrade
```

Note: If you want to be notified about new releases, please Watch (Releases only) the Lumispy repository on GitHub (requires a GitHub account).

1.2 Introduction

1.2.1 What is LumiSpy

LumiSpy is an open-source python library aimed at helping with the analysis of luminescence spectroscopy data – the development started mainly with photoluminescence (PL), cathodoluminescence (CL), electroluminescence (EL) and Raman spectroscopy in mind. Besides the standard continuous-excitation spectral data, the idea is to provide tools also for the analysis of time-resolved (transient) measurements. However, it may prove useful also for other optical measurements, such as absorption or transmission spectroscopy, scanning optical near field microscopy (SNOM), as well as fourier-transform infrared spectroscopy (FTIR).
**LumiSpy** is an extension to the python package **HyperSpy** that facilitates hyperspectral data analysis, i.e. maps or linescans where a spectrum is collected at each pixel. Or more general, multidimensional datasets that can be described as multidimensional arrays of a given signal.

Notable features that **HyperSpy** provides are:

- **base signal classes** for the handling of (multidimensional) spectral data,
- the necessary tools for loading **various data file formats**,
- **analytical tools** that exploit the multidimensionality of datasets,
- a user-friendly and powerful framework for **model fitting** that provides many standard functions and can be easily extended to custom ones,
- **machine learning** algorithms that can be useful, e.g. for denoising data,
- efficient handling of **big datasets**,
- **functions for data visualization** both to evaluate datasets during the analysis and provide interactive operation for certain functions, as well as for plotting of data,
- extracting subsets of data from multidimensional datasets via **regions of interest** and a powerful numpy-style indexing mechanism,
- handling of **non-uniform data axes** (introduced in the v1.7 release).

**LumiSpy** provides in particular:

- additional **Signal types** specifically for luminescence spectra and transients,
- transformation to **non-uniform signal axes** for use of other common units, such as eV (electron volt) and wavenumbers (Raman shift),
- additional **signal tools** such as data normalization and scaling,
- various **utility functions** useful in luminescence spectroscopy data analysis, such as joining multiple spectra along the signal axis, unit conversion, etc.

**LumiSpy** should facilitate an easy and reproducible analysis of single spectra or spectral images.

### 1.2.2 Signal types

As an extension to HyperSpy, LumiSpy provides several signal types extending the **base classes available in HyperSpy**. When the LumiSpy library is installed, these additional signal types are directly available to HyperSpy. To print all available specialised **hyperspy.signal.BaseSignal** subclasses installed in your system call the **hyperspy.utils.print_known_signal_types()** function:

```python
>>> import hyperspy.api as hs
>>> hs.print_known_signal_types()
```

The different subclasses are characterized by the **signal_type** metadata attribute. Some additional properties are summarized in the table below. Depending on the use case, certain functions will only be available for some signal types (or inheriting) signal types.
Table 1: LumiSpy subclasses and their basic attributes.

<table>
<thead>
<tr>
<th>BaseSignal subclass</th>
<th>signal_dimension</th>
<th>signal_type</th>
<th>dtype</th>
<th>aliases</th>
</tr>
</thead>
<tbody>
<tr>
<td>LumiSpectrum</td>
<td>1</td>
<td>Luminescence</td>
<td>real</td>
<td>LumiSpectrum, LuminescenceSpectrum</td>
</tr>
<tr>
<td>CLSpectrum</td>
<td>1</td>
<td>CL</td>
<td>real</td>
<td>CLSpectrum, cathodoluminescence</td>
</tr>
<tr>
<td>CLSEMSpectrum</td>
<td>1</td>
<td>CL_SEM</td>
<td>real</td>
<td>CLSEM, cathodoluminescence STEM</td>
</tr>
<tr>
<td>CLSTEMSpectrum</td>
<td>1</td>
<td>CL_STEM</td>
<td>real</td>
<td>CLSTEM, cathodoluminescence STEM</td>
</tr>
<tr>
<td>ELSpectrum</td>
<td>1</td>
<td>EL</td>
<td>real</td>
<td>ELSpectrum, electroluminescence</td>
</tr>
<tr>
<td>PLSpectrum</td>
<td>1</td>
<td>PL</td>
<td>real</td>
<td>PLSpectrum, photoluminescence</td>
</tr>
<tr>
<td>LumiTransient</td>
<td>1</td>
<td>Transient</td>
<td>real</td>
<td>TRLumi, TR luminescence, time-resolved luminescence</td>
</tr>
<tr>
<td>LumiTransientSpec</td>
<td>2</td>
<td>TransientSpec</td>
<td>real</td>
<td>TRLumiSpec, TR luminescence spectrum, time-resolved luminescence spectrum</td>
</tr>
</tbody>
</table>

The hierarchy of the LumiSpy signal types and their inheritance from HyperSpy is summarized in the following diagram:

```
    hyperspy.signal.BaseSignal
      ├── hyperspy._signals.signal1d.Signal1D
      │    ├── LumiSpectrum
      │    │    ├── CLSpectrum
      │    │    │    └── CLSEMSpectrum
      │    │    │    └── CLSTEMSpectrum
      │    │    └── ELSpectrum
      │    └── PLSpectrum
      └── LumiTransient
        └── hyperspy.signal.Signal2D
            └── LumiTransientSpectrum
```

1.2.3 Where are we heading?

LumiSpy is under active development, and as a user-driven project, we welcome contributions (see Contributing) to the code and documentation, but also bug reports and feature requests from any other users. Don’t hesitate to join the discussions!

Currently, we have implemented the base functionality that extends HyperSpy’s capabilities to additional signal classes. In the near future, the following functions should be developed:

- handling of transient (time-resolved) data,
- reading of common PL data formats,
- more dedicated analysis functionalities,
- ...

Chapter 1. Contents
1.3 Non-uniform signal axes

LumiSpy facilitates the use of non-uniform axes, where the points of the axis vector are not uniformly spaced. This situation occurs in particular when converting a wavelength scale to energy (eV) or wavenumbers (e.g. for Raman shifts).

The conversion of the signal axis can be performed using the functions to_eV(), to_invcm() and to_raman_shift() (alias for to_invcm_relative()). If the unit of the signal axis is set, the functions can handle wavelengths in either nm or µm.

Accepted parameters are inplace=True/False (default is True), which determines whether the current signal object is modified or a new one is created, and jacobian=True/False (default is True, see Jacobian transformation).

1.3.1 The energy axis

The transformation from wavelength \( \lambda \) to energy \( E \) is defined as \( E = \frac{hc}{\lambda} \). Taking into account the refractive index of air and doing a conversion from nm to eV, we get:

\[
E[\text{eV}] = \frac{10^9}{e} \frac{hc}{n_{\text{air}} \lambda[\text{nm}]},
\]

where \( h \) is the Planck constant, \( c \) is the speed of light, \( e \) is the elementary charge and \( n_{\text{air}} \) is the refractive index of air, see also [Pfueller].

```python
>>> s2 = s.to_eV(inplace=False)
>>> s.to_eV()
```

Note: The refractive index of air \( n_{\text{air}} \) is wavelength dependent. This dependence is taken into account by LumiSpy based on the analytical formula given by [Peck] valid from 185-1700 nm (outside of this range, the values of the refractive index at the edges of the range are used and a warning is raised).

1.3.2 The wavenumber axis/Raman shifts

The transformation from wavelength \( \lambda \) to wavenumber \( \tilde{\nu} \) (spatial frequency of the wave) is defined as \( \tilde{\nu} = 1/\lambda \). The wavenumber is usually given in units of cm\(^{-1}\).

When converting a signal to Raman shift, i.e. the shift in wavenumbers from the exciting laser wavelength, the laser wavelength has to be passed to the function using the parameter laser using the same units as for the original axis (e.g. 325 for nm or 0.325 for µm) unless it is contained in the LumiSpy metadata structure under Acquisition_instrument.Laser.wavelength.

TODO: Automatically read laser wavelength from metadata if given there.

```python
>>> s2 = s.to_invcm(inplace=False)
>>> s.to_invcm()
>>> s2 = s.to_raman_shift(inplace=False, laser=325)
>>> s.to_raman_shift(laser=325)
```

**1.3. Non-uniform signal axes**
1.3.3 Jacobian transformation

When transforming the signal axis, the signal intensity is automatically rescaled (Jacobian transformation), unless the `jacobian=False` option is given. Only converting the signal axis, and leaving the signal intensity unchanged, implies that the integral of the signal over the same interval would lead to different results depending on the quantity on the axis (see e.g. [Mooney] and [Wang]).

For the energy axis as example, if we require \( I(E)dE = I(\lambda)d\lambda \), then \( E = \frac{hc}{\lambda} \) implies

\[
I(E) = I(\lambda) \frac{d\lambda}{dE} = I(\lambda) \frac{hc}{dE} = -I(\lambda) \frac{hc}{E^2}
\]

The minus sign just reflects the different directions of integration in the wavelength and energy domains. The same argument holds for the conversion from wavelength to wavenumber (just without the additional prefactors in the equation). The renormalization in LumiSpy is defined such that the intensity is converted from counts/nm (or counts/\( \mu \)m) to counts/meV. The following figure illustrates the effect of the Jacobian transformation:

![Graph](image)

**Transformation of the variance**

Scaling the signal intensities implies that also the stored variance of the signal needs to be scaled accordingly. According to \( Var(aX) = a^2 Var(X) \), the variance has to be multiplied with the square of the Jacobian. This squared renormalization is automatically performed by LumiSpy if `jacobian=True`. In particular, homoscedastic (constant) noise will consequently become heteroscedastic (changing as a function of the signal axis vector). Therefore, if the `metadata.Signal.Noise_properties.variance` attribute is a constant, it is converted into a `hyperspy.signal.BaseSignal` object before the transformation.

See *Signal variance (noise)* for more general information on data variance in the context of model fitting and the HyperSpy documentation on setting the noise properties.

**Note:** If the Jacobian transformation is performed, the values of `metadata.Signal.Noise_properties.Variance_linear_model` are reset to their default values (`gain_factor=1`, `gain_offset=0` and `correlation_factor=1`). Should these values deviate from the defaults, make sure to run `hyperspy.signal.BaseSignal.estimate_poissonian_noise_variance()` prior to the transformation.
1.4 Signal tools

This section summarizes functions operating on the signal data. Besides those implemented in LumiSpy, it highlights functions from HyperSpy that are particularly useful for luminescence spectroscopy data.

1.4.1 Scaling and normalizing signal data

For comparative plotting or a detailed analysis, the intensity of spectra may need to be either scaled by the respective integration times or normalized. The luminescence signal classes provide these functionalities in the methods `scale_by_exposure()` and `normalize()`.

Both functions can operate directly on the signal (`inplace=True`), but as default a new signal is returned.

The scaling function can use the `integration_time` (unit: seconds) provided in the LumiSpy metadata structure (`metadata.Acqusition_instrument.Detector.integration_time`). Otherwise, the appropriate parameter has to be passed to the function.

```python
>>> scaled = s.scale_by_exposure(integration_time=0.5, inplace=True)
```

Normalization is performed for the pixel with maximum intensity. Alternatively, the parameter `pos` in calibrated units of the signal axis can be given to normalize the intensity at this position. Normalization may be convenient for plotting, but should usually not be performed on signals used as input for further analysis (therefore the default is `inplace=False`).

```python
>>> s.normalize(pos=450)
```

1.4.2 Peak positions and properties

Peak identification

HyperSpy provides functions to find the positions of maxima or minima in a dataset:

- `indexmax()` - return the index of the maximum value along a given axis.
- `indexmin()` - return the index of the minimum value along a given axis.
- `valuemax()` - return the position/coordinates of the maximum value along a given axis in calibrated units.
- `valuemin()` - return the position/coordinates of the minimum value along a given axis in calibrated units.

These functions take the `axis` keyword to define along which axis to perform the operation and return a new signal containing the result.

A much more powerful method to identify peaks is using the peak finding routine based on the downward zero-crossings of the first derivative of a signal: `find_peaks1D_ohaver()`. This function can find multiple peaks in a dataset and has a number of parameters for fine-tuning the sensitivity, etc.

All of these functions can be performed for a subset of the dataset:

```python
>>> peaks = s.find_peaks1D_ohaver()
>>> peaks = s.isig[100:-100].find_peaks1D_ohaver()
```
Peak Width

For asymmetric peaks, fitted functions may not provide an accurate description of the peak, in particular the peak width. The function `estimate_peak_width()` determines the width of a peak at a certain fraction of its maximum value. The default value `factor=0.5` returns the full width at half maximum (FWHM).

```python
>>> s.remove_background()
>>> width = s.estimate_peak_width(factor=0.3)
```

Calculating the centroid of a spectrum (centre of mass)

The function `centroid()` (based on the utility function `com()`) is an alternative to finding the position of the maximum intensity of a peak, useful in particular for non-symmetric peaks with pronounced shoulders. It finds the centroid (center of mass) of a peak in the spectrum from the signal axis units (or pixel number) and the intensity at each pixel value. It basically represents a “weighted average” of the peak as such:

\[
\text{com} = \frac{\sum x_i I_i}{\sum I_i},
\]

where \(x_i\) is the wavelength (or pixel number) at which the intensity of the spectrum \(I_i\) is measured.

This function also works for non-linear axes. For the `hyperspy.axes.FunctionalDataAxis`, the centroid is extrapolated based on the function used to create the non-uniform axis. For `hyperspy.axes.DataAxis`, a linear interpolation between the axes points at the center of mass is assumed, but this behaviour can be changed with the `kwargs` of `scipy.interpolate.interp1d` function.

```python
>>> s = lum.signals.LumiSpectrum([[1, 2, 3, 2, 1, 0]*2]*3)
>>> s
<LumiSpectrum, title: , dimensions: (2|6|3)

>>> ax = s.axes_manager.signal_axes[0]
>>> ax.offset = 200
>>> ax.scale = 100

>>> com = s.centroid()
>>> com
<Signal2D, title: Centroid map, dimensions: (2|3)

400.0
```

Note: This function only works for a single peak. If you have multiple peaks, slice the signal beforehand or use the slice parameter (which follows the `s.isig[:]` convention).

Note: The Jacobian transformation may affect the shape, in particular of broader peaks. It is therefore highly recommended to convert luminescence spectra from wavelength to the energy axis prior to determining the centroid to determine the true emission energy. See e.g. [Wang] and [Mooney].
1.4.3 Signal statistics and analytical operations

Standard statistical operations can be performed on the data or a subset of the data, notably these include `max()`, `min()`, `sum()`, `mean()`, `std()`, and `var()`. Variations of all these functions exist that ignore missing values (NaN) if present, e.g. `nanmax()`.

Integration along a specified signal axis is performed using the function `integrate1D()`.

The numerical derivative of a signal can be calculated using the function `derivative()`, while the n-th order discrete difference can be calculated using `diff()`.

These functions take the `axis` keyword to define along which axis to perform the operation and return a new signal containing the result:

```python
>>> area = s.integrate1D(axis=0)
```

1.4.4 Replacing negative data values

Log-scale plotting fails in the presence of negative values in the dataset (e.g. introduced after background removal). In this case, the utility function `remove_negative()` replaces all negative values in the data array by a basevalue (default basevalue=1). The default operational mode is `inplace=False` (a new signal object is returned).

```python
>>> s.remove_negative(0.1)
```

1.5 Fitting luminescence data

LumiSpy is compatible with HyperSpy model fitting. It can fit using both uniform and non-uniform axes (e.g. energy scale). A general introduction can be found in the HyperSpy user guide.

A detailed example is given in the `Fitting_tutorial` in the HyperSpy demos repository. See also the LumiSpy demo notebooks for examples of data fitting.

**Note:** The *Jacobian transformation* may affect the shape, in particular of broader peaks. It is therefore highly recommended to convert luminescence spectra from wavelength to the *energy axis* prior to any fitting to obtain the true emission energy. See e.g. [Wang] and [Mooney].

TODO: Show how to extract the *modeled signal* with all/one component.

1.5.1 Signal variance (noise)

TODO: Documentation on variance handling in the context of fitting, in particular using `estimate_poissonian_noise_variance()`

For a detailed discussion, see [Tappy]
1.6 Utility functions

This section summarizes various useful functions implemented in LumiSpy.

1.6.1 Join spectra

In case several spectra (or spectral images) where subsequently recorded for different, but overlapping spectral windows, LumiSpy provides a utility `join_spectra()` to merge these into a single spectrum. The main argument is a list of two or more spectral objects. Spectra are joined at the centre of the overlapping range along the signal axis. To avoid steps in the intensity, several parameters (see docstring: `join_spectra()`) allow to tune the scaling of the later signals with respect to the previous ones. By default, the scaling parameter is determined as average ratio between the two signals in the range of +/- 50 pixels around the centre of the overlapping region.

```python
>>> import lumispy as lum
>>> s = lum.join_spectra((s1,s2))
```

1.6.2 Utilities for spectral maps

The function `crop_edges()` removes the specified number of pixels from all four edges of a spectral map. It is a convenience wrapper for the `inav` method in HyperSpy.

```python
>>> s.crop_edges(crop_px=2)
```

[TODO: add possibility to crop different amounts of pixels on different sides]

1.6.3 Unit conversion

For convenience, LumiSpy provides functions that convert between different units commonly used for the signal axis. Namely,

- `nm2eV()`
- `eV2nm()`
- `nm2invcm()`
- `invcm2nm()`

For the energy axis, the conversion uses the wavelength-dependent refractive index of air.

1.6.4 Solving the grating equation

The function `solve_grating_equation()` follows the conventions described in the tutorial from Horiba Scientific.
1.7 LumiSpy metadata structure

LumiSpy extends the HyperSpy metadata structure with conventions for metadata specific to its signal types. Refer to the HyperSpy metadata documentation for general metadata fields.

The metadata of any signal objects is stored in the metadata attribute, which has a tree structure. By convention, the node labels are capitalized and the ones for leaves are not capitalized. When a leaf contains a quantity that is not dimensionless, the units can be given in an extra leaf with the same label followed by the _units suffix.

Besides directly accessing the metadata tree structure, e.g. s.metadata.Signal.signal_type, the HyperSpy methods set_item(), has_item() and get_item() can be used to add to, search for and read from items in the metadata tree, respectively.

The luminescence specific metadata structure is represented in the following tree diagram. The default units are given in parentheses. Details about the leaves can be found in the following sections of this chapter. Note that not all types of leaves will apply to every type of measurement. For example, while parallel acquisition with a CCD is characterized by the central_wavelength, a serial acquisition with a PMT will require a start_wavelength and a step_size.

metadata
  ┐ General
  │   └ # see HyperSpy
  ┌ Sample
  │   └ # see HyperSpy
  ┌ Signal
  │   │ signal_type
  │   │ quantity
  │   │   # otherwise see HyperSpy
  │ Acquisition_instrument
  │   └ Laser / SEM / TEM
  │       │ laser_type
  │       │ model
  │       │ wavelength (nm)
  │       │ power (mW)
  │       │ objective_magnification
  │       │ Filter
  │       │       │ filter_type
  │       │       │ position
  │       │       │ optical_density
  │       │       │ cut_on_wavelength (nm)
  │       │       │ cut_off_wavelength (nm)
  │       │   # for SEM/TEM see HyperSpy
  │       └ Spectrometer
  │           │ model
  │           │ acquisition_mode
  │           │ entrance_slit_width (mm)
  │           │ exit_slit_width (mm)
  │           │ central_wavelength (nm)
  │           │ start_wavelength (nm)
  │           │ step_size (nm)
  │           │ Grating
  │           │       │ groove_density (grooves/mm)
  │           │       │ blazing_angle (°)
  │           │       │ blazing_wavelength (nm)
  │           │ Filter

(continues on next page)
1.7.1 General

See HyperSpy-Metadata-General.

1.7.2 Sample

See HyperSpy-Metadata-Sample.

1.7.3 Signal

signal_type
type: string

String that describes the type of signal. The LumiSpy specific signal classes are summarized under Signal types.

quantity
type: string

The name of the quantity of the “intensity axis” with the units in round brackets if required, for example ‘Intensity (counts/s)’.

See HyperSpy-Metadata-Signal for additional fields.
1.7.4 Acquisition Instrument

1.7.5 Laser / SEM / TEM

For SEM or TEM see HyperSpy-Metadata-SEM or HyperSpy-Metadata-TEM.

Laser

laser_type
type: string
The type of laser used, e.g. ‘HeCd’.

model
type: string
Model of the laser (branding by manufacturer).

wavelength
type: float
Emission wavelength of the exciting laser in nm.

power
type: float
Measured power of the excitation laser in mW.

magnification
type: int
Magnification of the microscope objective used to focus the beam to the sample.

Filter

Information about additional filters entered into the lightpath before the sample. In case multiple filters are used, they should be numbered Filter_1, etc.

filter_type
type: string
Type of filter (e.g. ‘optical density’, ‘short pass’, ‘long pass’, ‘bandpass’, ‘color’).

position
type: string
Position in the beam (e.g. ‘excitation’ vs. ‘detection’ in case of optical excitation).

optical_density
type: float
Optical density in case of an intensity filter.

cut_on_wavelength
type: float
Cut on wavelength in nm in case of a long-pass or bandpass filter.

cut_off_wavelength
type: float
Cut off wavelength in nm in case of a short-pass or bandpass filter.
1.7.6 Spectrometer

Contains information about the spectrometer, configuration and grating used for the measurement. In case multiple spectrometers are connected in series, they should be numbered Spectrometer_1, etc.

**model**
- type: string
  Model of the spectrometer (branding by manufacturer).

**acquisition_mode**
- type: string
  Acquisition mode (e.g. ‘Parallel dispersive’, versus ‘Serial dispersive’).

**entrance_slit_width**
- type: float
  Width of the entrance slit in mm.

**exit_slit_width**
- type: float
  Width of the exit slit (serial acquisition) in mm.

**central_wavelength**
- type: float
  Central wavelength during acquisition (parallel acquisition).

**start_wavelength**
- type: float
  Start wavelength in nm (serial acquisition).

**step_size**
- type: float
  Step size in nm (serial acquisition).

**Grating**

Information of the dispersion grating employed in the measurement.

**groove_density**
- type: int
  Density of lines on the grating in grooves/mm.

**blazing_angle**
- type: int
  Angle in degree (°) that the grating is blazed at.

**blazing_wavelength**
- type: int
  Wavelength that the grating blaze is optimized for in nm.
Filter

Information about additional filters entered into the lightpath after the sample. In case multiple filters are used, they should be numbered Filter_1, etc. See Filter above for details on items that may potentially be included.

1.7.7 Detector

Contains information about the detector used to acquire the signal. Contained leaves will differ depending on the type of detector.

detector_type
type: string
The type of detector used to acquire the signal (CCD, PMT, StreakCamera, TCSPD).

model
type: string
The model of the used detector.

frames
type: int
Number of frames that are summed to yield the total integration time.

integration_time (s)
type: float
Time over which the signal is integrated. In case multiple frames are summed, it is the total exposure time. In case of serial acquisition, it is the dwell time per data point.

saturation_fraction
type: float
Fraction of the signal intensity compared with the saturation threshold of the CCD.

binning
type: tuple of int
A tuple that describes the binning of a parallel detector such a CCD on readout in x and y directions.

processing
type: string
Information about automatic processing performed on the data, e.g. ‘dark subtracted’.

sensor_roi
type: tuple of int
Tuple of length 2 or 4 that specifies range of pixels on a detector that are read out: (offset x, offset y, size x, size y) for a 2D array detector and (offset, size) for a 1D line detector.

pixel_size
type: float or tuple of float
Size of a pixel in µm. Tuple of length 2 (width, height), when the pixel is not square.
1.7.8 Spectral_image

Contains information about mapping parameters, such as step size, drift correction, etc.

mode
type: string
Mode of the spectrum image acquisition such as ‘Map’ or ‘Linescan’.

drift_correction_periodicity
type: int/float
Periodicity of the drift correction in specified units (standard s).

drift_correction_units
type: string
Units of the drift correction such as ‘s’, ‘px’, ‘rows’.

1.8 Bibliography

1.9 lumispy

1.9.1 lumispy package

Subpackages

lumispy.signals package

Submodules

lumispy.signals.cl_spectrum module

Signal class for cathodoluminescence spectral data

class lumispy.signals.cl_spectrum.CLSEMSpectrum(*args, **kwargs)
Bases: CLSpectrum

1D scanning electron microscopy cathodoluminescence signal class.

correct_grating_shift(cal_factor_x_axis, corr_factor_grating, sem_magnification, **kwargs)
Applies shift caused by the grating offset wrt the scanning centre. Authorship: Gunnar Kusch (gk419@cam.ac.uk)

Parameters

- cal_factor_x_axis – The navigation correction factor.
- corr_factor_grating – The grating correction factor.
- sem_magnification – The SEM (real) magnification value. For the Attolight original metadata, take the SEM.Real_Magnification value
class lumispy.signals.cl_spectrum.CLSTEMSpectrum(*args, **kwargs)

Bases: CLSpectrum

1D scanning transmission electron microscopy cathodoluminescence signal class.

class lumispy.signals.cl_spectrum.CLSpectrum(*args, **kwargs)

Bases: LumiSpectrum

General 1D cathodoluminescence signal class.

_make_signal_mask(luminescence_roi)

Creates a mask from the peak position and peak widths of the luminescence spectrum.

Parameters

- luminescence_roi (array) – In the form of an array of pairwise elements \([\text{peak1}_x, \text{peak1}_\text{width}], [\text{peak2}_x, \text{peak2}_\text{width}]....\).

Returns

A signal_mask.

Return type

array

remove_spikes(threshold='auto', show_diagnosis_histogram=False, inplace=False, luminescence_roi=None, signal_mask=None, add_noise=False, navigation_mask=None, interactive=False, **kwargs)

HyperSpy-based spike removal tool adapted to LumiSpy to run non-interactively and without noise addition by default. Graphical interface to remove spikes from EELS spectra or luminescence data. If non-interactive, it removes all spikes and returns a ~hyperspy.signals._signal_tools.SpikesRemoval object.

Parameters

- **signal_mask (bool array) –** Restricts the operation to the signal locations not marked as True (masked).
- **navigation_mask (bool array) –** Restricts the operation to the navigation locations not marked as True (masked).
- **threshold (‘auto’ or int) –** If int, set the threshold value used for detecting the spikes. If auto, determine the threshold value as being the first zero value in the histogram obtained from the spikes_diagnosis() method.
- **interactive (bool) –** If True, remove the spikes using the graphical user interface. If False, remove all the spikes automatically, which can introduce artefacts if used with signal containing peak-like features. However, this can be mitigated by using the signal_mask argument to mask the signal of interest.
- **display (bool) –** If True, display the user interface widgets. If False, return the widgets container in a dictionary, usually for customisation or testing.
- **toolkit (str, iterable of strings or None) –** If None (default), all available widgets are displayed or returned. If string, only the widgets of the selected toolkit are displayed if available. If an iterable of toolkit strings, the widgets of all listed toolkits are displayed or returned.
- ****kwargs (dict) –

Keyword arguments pass to

SpikesRemoval()

See also:

spikes_diagnosis()
Parameters

- `show_diagnosis_histogram (bool)` – Plot or not the derivative histogram to show the magnitude of the spikes present.
- `inplace (bool)` – If False, a new signal object is created and returned. If True, the original signal object is modified.
- `luminescence_roi (array)` – The peak position and peak widths of the peaks in the luminescence spectrum. In the form of an array of pairwise elements `[[peak1_x, peak1_width], [peak2_x, peak2_width], ...]` in the units of the signal axis. It creates a signal_mask protecting the peak regions. To be used instead of `signal_mask`.

Returns

Depends on inplace, returns or overwrites the CLSpectrum after spike removal.

Return type

None or `CL Spectrum`

class lumispy.signals.cl_spectrum.LazyCLSEM Spectrum(*args, **kwargs)
    Bases: LazySignal, CLSEM Spectrum

Lazy 1D scanning electron microscopy cathodoluminescence signal class.

class lumispy.signals.cl_spectrum.LazyCLSTEM Spectrum(*args, **kwargs)
    Bases: LazySignal, CLSTEM Spectrum

Lazy 1D scanning transmission electron microscopy cathodoluminescence signal class.

class lumispy.signals.cl_spectrum.LazyCLSpectrum(*args, **kwargs)
    Bases: LazySignal, CLSpectrum

General lazy 1D cathodoluminescence signal class.

lumispy.signals.common_luminescence module

Signal class for luminescence data (BaseSignal class)

class lumispy.signals.common_luminescence.CommonLumi
    Bases: object

General luminescence signal class (dimensionless)

crop_edges(crop_px)
    Crop the amount of pixels from the four edges of the scanning region, from out the edges inwards.

Parameters

crop_px (int) – Amount of pixels to be cropped on each side individually.

Returns

signal_cropped – A smaller cropped CL signal object. If inplace is True, the original object is modified and no LumiSpectrum is returned.

Return type

CommonLuminescence
normalize(pos=nan, element_wise=False, inplace=False)

Normalizes data to value at pos along signal axis, defaults to maximum value.

Can be helpful for e.g. plotting, but does not make sense to use on signals that will be used as input for further calculations!

Parameters

• pos (float, int) – If ‘nan’ (default), spectra are normalized to the maximum. If float, position along signal axis in calibrated units at which to normalize the spectra. If int, index along signal axis at which to normalize the spectra.

• element_wise (boolean) – If False (default), a spectrum image is normalized by a common factor. If True, each spectrum is normalized individually.

• inplace (boolean) – If False (default), a new signal object is created and returned. If True, the operation is performed on the existing signal object.

Notes

Sets metadata.Signal.normalized to True. If metadata.Signal.quantity contains the word ‘Intensity’, replaces this field with ‘Normalized intensity’.

remove_negative(basevalue=1, inplace=False)

Sets all negative values to ‘basevalue’, e.g. for logarithmic scale plots.

Parameters

• basevalue (float) – Value by which negative values are replaced (default = 1).

• inplace (boolean) – If False (default), a new signal object is created and returned. Otherwise, the operation is performed on the existing signal object.

Notes

Sets metadata.Signal.negative_removed to True.

scale_by_exposure(integration_time=None, inplace=False, **kwargs)

Scale data in spectrum by integration time / exposure, (e.g. convert counts to counts/s).

Parameters

• integration_time (float) – Integration time (exposure) in s. If not given, the function tries to use the ‘metadata.Acqusition_instrument.Detector.integration_time’ field or alternatively find any ‘integration_time’, ‘exposure’ or ‘dwell_time’ fields in the original_metadata.

• inplace (boolean) – If False (default), a new signal object is created and returned. If True, the operation is performed on the existing signal object.
Notes

Sets metadata.Signal.scaled to True. If intensity units is ‘counts’, replaces them by ‘counts/s’.

Deprecated since version 0.2: The exposure argument was renamed integration_time, and it will be removed in LumiSpy 1.0.

lumispy.signals.common_transient module

Signal class for transient data (BaseSignal class)

class lumispy.signals.common_transient.CommonTransient
    Bases: object
    General transient signal class (dimensionless)

lumispy.signals.el_spectrum module

Signal class for electroluminescence spectral data

class lumispy.signals.el_spectrum.ELSpectrum(*args, **kwargs)
    Bases: LumiSpectrum
    General 1D electroluminescence signal class

class lumispy.signals.el_spectrum.LazyELSpectrum(*args, **kwargs)
    Bases: LazySignal, ELSpectrum
    General lazy 1D electroluminescence signal class

lumispy.signals.luminescence_spectrum module

Signal class for Luminescence spectral data (1D).

class lumispy.signals.luminescence_spectrum.LazyLumiSpectrum(*args, **kwargs)
    Bases: LazySignal, LumiSpectrum
    General lazy 1D luminescence signal class.

class lumispy.signals.luminescence_spectrum.LumiSpectrum(*args, **kwargs)
    Bases: Signal1D, CommonLumi
    General 1D luminescence signal class.

    _convert_data(newaxis, factor, inplace, jacobian, data2, var2)
        Utility function to perform the data and variance conversion for signal unit transformations.

    _reset_variance_linear_model()
        Resets the variance linear model parameters to their default values, as they are not applicable any longer after a Jacobian transformation.

    centroid(signal_range=None, **kwargs)
        Finds the centroid (center of mass) of a peak in the spectrum from the wavelength (or pixel number) and the intensity at each pixel value. It basically represents a “weighted average” of the peak.
Notes

This function only works for a single peak. If you have multiple peaks, slice the signal beforehand or use the signal_range parameter.

TODO: Implement this function for multiple peaks (with the npeaks parameter) by finding the top 2 peaks from mean spectrum and then returning a signal with 2 com.

Parameters

- **signal_range** *(tuple of ints or floats, optional)* – A tuple representing the indices of the signal axis (start index, end index) where the peak is located. If the tuple contains int, it slices on index. If the tuple contains float, it slices on signal units (default HyperSpy s.inav[:]) functionality.

- **kwargs** *(dictionary)* – For the scipy.interpolate.interp1d function.

Returns

- **signal** – Signal object containing the center of mass for every pixel. Depending on the dimensionality the type is Signal2D or a BaseSignal (for single spectrum).

Return type

Signal2D, BaseSignal

px_to_nm_grating_solver

- **gamma_deg** *(float)* – Inclination angle between the focal plane and the centre of the grating (found experimentally from calibration). In degree.

- **deviation_angle_deg** *(float)* – Also known as included angle. It is defined as the difference between angle of diffraction ($\beta$) and angle of incidence ($\alpha$). Given by manufacturer specsheet. In degree.

- **focal_length_mm** *(float)* – Given by manufacturer specsheet. In mm.

- **ccd_width_mm** *(float)* – The width of the CDD. Given by manufacturer specsheet. In mm.

- **grating_central_wavelength_nm** *(float)* – Wavelength at the centre of the grating, where exit slit is placed. In nm.

- **grating_density_gr_mm** *(int)* – Grating density in gratings per mm.

- **inplace** *(bool)* – If False, it returns a new object with the transformation. If True, the original object is transformed, returning no object.

Returns

- **signal** – A signal with calibrated wavelength units.

Return type

LumiSpectrum
Examples

```python
>>> s = LumiSpectrum(np.ones(20),)
>>> s.px_to_nm_grating_solver(*params, inplace=True)
>>> s.axes_manager.signal_axes[0].units == 'nm'
```

`remove_background_from_file(…)`

Subtract the background to the signal in all navigation axes. If no background file is passed as argument, the `remove_background()` from HyperSpy is called with the GUI.

**Parameters**

- **background** *(array shape (2, n) or Signal1D)* – An array containing the background x-axis and the intensity values [[xs],[ys]] or a Signal1D object. If the x-axis values do not match the signal_axes, then interpolation is done before subtraction. If only the intensity values are provided, [ys], the functions assumes no interpolation needed.
- **inplace** *(boolean)* – If False, it returns a new object with the transformation. If True, the original object is transformed, returning no object.

**Returns**

- **signal** – A background subtracted signal.

**Return type**

`LumiSpectrum`

**Notes**

This function does not work with non-uniform axes.

`savetxt(…)`

Writes luminescence spectrum object to simple text file.

Writes single spectra to a two-column data file with signal axis as X and data as Y. Writes linescan data to file with signal axis as first row and navigation axis as first column (flipped if `transpose=True`).

**Parameters**

- **filename** *(string)* –
- **fmt** *(str or sequence of strs, optional)* – A single or sequence of format strings. Default is '%.5f'.
- **delimiter** *(str, optional)* – String or character separating columns. Default is ','. 
- **axes** *(bool, optional)* – If True (default), include axes in saved file. If False, save the data array only.
- **transpose** *(bool, optional)* – If True, transpose data array and exchange axes. Default is false. Ignored for single spectra.
- ****kwargs** – Takes any additional arguments of `numpy.savetxt`, e.g. `newline`, `header`, `footer`, `comments`, or `encoding`.

**See also:**

`numpy.savetxt`
Examples

```python
>>> import lumispy as lum
>>> import numpy as np
...
>>> # Spectrum:
>>> S = lum.signals.LumiSpectrum(np.arange(5))
>>> S.savetxt('spectrum.txt')
0.00000 0.00000
1.00000 1.00000
2.00000 2.00000
3.00000 3.00000
4.00000 4.00000
...
>>> # Linescan:
>>> L = lum.signals.LumiSpectrum(np.arange(25).reshape((5,5)))
>>> L.savetxt('linescan.txt')
0.00000 0.00000 1.00000 2.00000 3.00000 4.00000
0.00000 0.00000 5.00000 10.00000 15.00000 20.00000
2.00000 2.00000 7.00000 12.00000 17.00000 22.00000
3.00000 3.00000 8.00000 13.00000 18.00000 23.00000
to_array(axes=True, transpose=False)
```

**to_array** *(axes=True, transpose=False)*

Returns luminescence spectrum object as numpy array (optionally including the axes).

Returns single spectra as two-column array. Returns linescan data as array with signal axis as first row and navigation axis as first column (flipped if transpose=True).

**Parameters**

- **axes** *(bool, optional)* – If True (default), include axes in array. If False, return the data array only.
- **transpose** *(bool, optional)* – If True, transpose data array and exchange axes. Default is false. Ignored for single spectra.
- ****kwargs **– Takes any additional arguments of numpy.loadtxt, e.g. newline header, footer, comments, or encoding.

**Notes**

The output of this function can be used to convert a signal object to a pandas dataframe, e.g. using `df = pd.DataFrame(S.to_array())`. 
Examples

```python
>>> import lumispy as lum
>>> import numpy as np
...
>>> # Spectrum:
>>> S = lum.signals.LumiSpectrum(np.arange(5))
>>> S.to_array()
array([[0., 0.],
       [1., 1.],
       [2., 2.],
       [3., 3.],
       [4., 4.]])
...
>>> # Linescan:
>>> L = lum.signals.LumiSpectrum(np.arange(25).reshape((5,5)))
>>> L.to_array()
array([[ 0.,  0.,  1.,  2.,  3.,  4.],
       [ 0.,  0.,  1.,  2.,  3.,  4.],
       [ 1.,  5.,  6.,  7.,  8.,  9.],
       [ 2., 10., 11., 12., 13., 14.],
       [ 3., 15., 16., 17., 18., 19.],
       [ 4., 20., 21., 22., 23., 24.]])
```

to_eV(inplace=True, jacobian=True)

Converts signal axis of 1D signal to non-linear energy axis (eV) using wavelength dependent refractive index of air. Assumes wavelength in units of nm unless the axis units are specifically set to µm.

The intensity is converted from counts/nm (counts/µm) to counts/meV by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013), doi:10.1021/jz401508t, which ensures that integrated signals are correct also in the energy domain. If the variance of the signal is known, i.e. `metadata.Signal.Noise_properties.variance` is a signal representing the variance, a squared renormalization of the variance is performed. Note that if the variance is a number (not a signal instance), it is converted to a signal if the Jacobian transformation is performed.

Parameters

- **inplace** (boolean) – If `False`, a new signal object is created and returned. Otherwise (default) the operation is performed on the existing signal object.
- **jacobian** (boolean) – The default is to do the Jacobian transformation (recommended at least for luminescence signals), but the transformation can be suppressed by setting this option to `False`.

Examples

```python
>>> import numpy as np
>>> from lumispy import LumiSpectrum
>>> S1 = LumiSpectrum(np.ones(20), DataAxis(axis = np.arange(200,400,10)), )
>>> S1.to_eV()
```

to_invcm(inplace=True, jacobian=True)

Converts signal axis of 1D signal to non-linear wavenumber axis (cm^-1). Assumes wavelength in units of nm unless the axis units are specifically set to µm.
The intensity is converted from counts/nm (counts/µm) to counts/cm^-1 by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013), doi:10.1021/jz401508t, which ensures that integrated signals are correct also in the wavenumber domain. If the variance of the signal is known, i.e. metadata.Signal.Noise_properties.variance is a signal representing the variance, a squared renormalization of the variance is performed. Note that if the variance is a number (not a signal instance), it is converted to a signal if the Jacobian transformation is performed.

**Parameters**

- **inplace** *(boolean)* – If *False*, a new signal object is created and returned. Otherwise (default) the operation is performed on the existing signal object.

- **jacobian** *(boolean)* – The default is to do the Jacobian transformation (recommended at least for luminescence signals), but the transformation can be suppressed by setting this option to *False*.

**Examples**

```python
>>> import numpy as np
>>> from lumispy import LumiSpectrum
>>> S1 = LumiSpectrum(np.ones(20), DataAxis(axis = np.arange(200,400,10)), )
>>> S1.to_invcm()
```

to_invcm_relative *(laser=None, inplace=True, jacobian=False)*

Converts signal axis of 1D signal to non-linear wavenumber axis (cm^-1) relative to the exciting laser wavelength (Raman/Stokes shift). Assumes wavelength in units of nm unless the axis units are specifically set to µm.

The intensity is converted from counts/nm (counts/µm) to counts/cm^-1 by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013), doi:10.1021/jz401508t, which ensures that integrated signals are correct also in the wavenumber domain. If the variance of the signal is known, i.e. metadata.Signal.Noise_properties.variance is a signal representing the variance, a squared renormalization of the variance is performed. Note that if the variance is a number (not a signal instance), it is converted to a signal if the Jacobian transformation is performed.

**Parameters**

- **inplace** *(boolean)* – If *False*, a new signal object is created and returned. Otherwise (default) the operation is performed on the existing signal object.

- **laser** *(float or None)* – Laser wavelength in the same units as the signal axis. If None (default), checks if it is stored in metadata.Acquisition_instrument.Laser.wavelength.

- **jacobian** *(boolean)* – The default is not to do the Jacobian transformation for Raman shifts, but the transformation can be activated by setting this option to *True*.

**Examples**

```python
>>> import numpy as np
>>> from lumispy import LumiSpectrum
>>> S1 = LumiSpectrum(np.ones(20), DataAxis(axis = np.arange(200,400,10)), )
>>> S1.to_invcm(laser=325)
```
to_raman_shift(laser=None, inplace=True, jacobian=False)

Converts signal axis of 1D signal to non-linear wavenumber axis (cm^{-1}) relative to the exciting laser wavelength (Raman/Stokes shift). Assumes wavelength in units of nm unless the axis units are specifically set to µm.

The intensity is converted from counts/nm (counts/µm) to counts/cm^{-1} by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013), doi:10.1021/jz401508t, which ensures that integrated signals are correct also in the wavenumber domain. If the variance of the signal is known, i.e. metadata.Signal.Noise_properties.variance is a signal representing the variance, a squared renormalization of the variance is performed. Note that if the variance is a number (not a signal instance), it is converted to a signal if the Jacobian transformation is performed.

Parameters

- **inplace** (boolean) – If False, a new signal object is created and returned. Otherwise (default) the operation is performed on the existing signal object.
- **laser** (float or None) – Laser wavelength in the same units as the signal axis. If None (default), checks if it is stored in metadata.Acquisition_instrument.Laser.wavelength.
- **jacobian** (boolean) – The default is not to do the Jacobian transformation for Raman shifts, but the transformation can be activated by setting this option to True.

Examples

```python
>>> import numpy as np
>>> from lumispy import LumiSpectrum
>>> S1 = LumiSpectrum(np.ones(20), DataAxis(axis = np.arange(200,400,10)),)
>>> S1.to_invcm(laser=325)
```

lumispy.signals.luminescence_transient module

Signal class for luminescence transient data (1D)

class lumispy.signals.luminescence_transient.LazyLumiTransient(*args, **kwargs)

Bases: LazySignal, LumiTransient

General lazy 1D luminescence signal class (transient/time resolved)

class lumispy.signals.luminescence_transient.LumiTransient(*args, **kwargs)

Bases: Signal1D, CommonTransient

General 1D luminescence signal class (transient/time resolved)

lumispy.signals.luminescence_transientspec module

Signal class for luminescence transient data (2D)

class lumispy.signals.luminescence_transientspec.LazyLumiTransientSpectrum(*args, **kwargs)

Bases: LazySignal, LumiTransientSpectrum

General lazy 2D luminescence signal class (transient/time resolved)
class lumispy.signals.luminescence_transient_spec.LumiTransientSpectrum(*args, **kwargs)

Bases: Signal2D, CommonLumi, CommonTransient

General 2D luminescence signal class (transient/time resolved)

lumispy.signals.pl_spectrum module

Signal class for Photoluminescence spectral data

class lumispy.signals.pl_spectrum.LazyPLSpectrum(*args, **kwargs)

Bases: LazySignal, PLSpectrum

General lazy 1D photoluminescence signal class

class lumispy.signals.pl_spectrum.PLSpectrum(*args, **kwargs)

Bases: LumiSpectrum

General 1D photoluminescence signal class

Module contents

lumispy.utils package

Submodules

lumispy.utils.axes module

lumispy.utils.axes._n_air(x)


lumispy.utils.axes.axis2eV(ax0)

Converts given signal axis to energy scale (eV) using wavelength dependent refractive index of air. Assumes wavelength in units of nm unless the axis units are specifically set to µm.

lumispy.utils.axes.axis2invcm(ax0)

Converts given signal axis to wavenumber scale (cm⁻¹). Assumes wavelength in units of nm unless the axis units are specifically set to µm.

lumispy.utils.axes.data2eV(data, factor, evaxis, ax0)

The intensity is converted from counts/nm (counts/µm) to counts/meV by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013). Ensures that integrated signals are still correct.

lumispy.utils.axes.data2invcm(data, factor, invcmaxis, ax0=None)

The intensity is converted from counts/nm (counts/µm) to counts/cm⁻¹ by doing a Jacobian transformation, see e.g. Mooney and Kambhampati, J. Phys. Chem. Lett. 4, 3316 (2013). Ensures that integrated signals are still correct.

lumispy.utils.axes.eV2nm(x)

Converts energy (eV) to wavelength (nm) using wavelength-dependent refractive index of air.

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lumispy.utils.axes.invcm2nm(x)
    Converts wavenumber (cm\(^{-1}\)) to wavelength (nm).

lumispy.utils.axes.join_spectra(S, r=50, scale=True, average=False, kind='slinear')
    Takes list of Signal1D objects and returns a single object with all spectra joined. Joins spectra at the center of the overlapping range. Scales spectra by a factor determined as average over the range center -/+ r pixels. Works both for uniform and non-uniform axes (FunctionalDataAxis is converted into a non-uniform DataAxis).

    Parameters
    - S (list of Signal1D objects (with overlapping signal axes)) --
    - r (int, optional) – Number of pixels left/right of center (default 50) defining the range over which to determine the scaling factor, has to be less than half of the overlapping pixels. Change the size of r or use average=True if the function induces a step in the intensity.
    - scale (boolean, optional) – If True (default), the later spectra in the list are scaled by a factor determined over center -/+ r pixels. If False, spectra are joined without scaling, which will likely induce a step unless average=True.
    - average (boolean, optional) – If True, the contribution of the two signals is continuously graded within the range defined by r instead of joining at the center of the range (default).
    - kind (str, optional) – Interpolation method (default ‘slinear’) to use when joining signals with a uniform signal axes. See scipy.interpolate.interp1d for options.

    Returns
    - A new Signal1D object containing the joined spectra (properties are copied from first spectrum).

    Examples
    >>> s1 = hs.signals.Signal1D(np.ones(32))
    >>> s2 = hs.signals.Signal1D(np.ones(32)*2)
    >>> s2.axes_manager.signal_axes[0].offset = 25
    >>> lum.join_spectra([s1,s2],r=2)
<Signal1D, title: , dimensions: (|57)>
• **deviation_angle_deg** *(float)* – Also known as included angle. It is defined as the difference between angle of diffraction ($\beta$) and angle of incidence ($\alpha$). Given by manufacturer specsheet. In degree.

• **focal_length_mm** *(float)* – Given by manufacturer specsheet. In mm.

• **ccd_width_mm** *(float)* – The width of the CDD. Given by manufacturer specsheet. In mm.

• **grating_central_wavelength_nm** *(float)* – Wavelength at the centre of the grating, where exit slit is placed. In nm.

• **grating_density_gr_mm** *(int)* – Grating density in gratings per mm.

Returns

axis – HyperSpy axis object.

Return type

hyperspy.axis

lumispy.utils.axes.var2eV(variance, factor, evaxis, ax0)

The variance is converted doing a squared Jacobian renormalization to match with the transformation of the data.

lumispy.utils.axes.var2invcm(variance, factor, invcmaxis, ax0=None)

The variance is converted doing a squared Jacobian renormalization to match with the transformation of the data.

lumispy.utils.io module

lumispy.utils.io.savetxt(S, filename, fmt='%.5f', delimiter='\t', axes=True, transpose=False, **kwargs)

Writes signal object to simple text file.

Writes single spectra to a two-column data file with signal axis as X and data as Y. Writes linescan data to file with signal axis as first row and navigation axis as first column (flipped if transpose=True). Writes image to file with the navigation axes as first column and first row. Writes 2D data (e.g. map of a fit parameter value) to file with the signal axes as first column and first row.

Parameters

• filename *(string)* –

• fmt *(str or sequence of strs, optional)* – A single or sequence of format strings. Default is '%.5f'.

• delimiter *(str, optional)* – String or character separating columns. Default is ' \t '.

• axes *(bool, optional)* – If True (default), include axes in saved file. If False, save the data array only.

• transpose *(bool, optional)* – If True, transpose data array and exchange axes. Default is false. Ignored for single spectra.

• **kwargs* – Takes any additional arguments of numpy.loadtxt, e.g. newline header, footer, comments, or encoding.

See also:

numpy.savetxt
Examples

```python
>>> import lumispy as lum
>>> import numpy as np
...
>>> # Spectrum:
>>> S = lum.signals.LumiSpectrum(np.arange(5))
>>> lum.savetxt(S, 'spectrum.txt')
0.00000  0.00000
1.00000  1.00000
2.00000  2.00000
3.00000  3.00000
4.00000  4.00000
...
>>> # Linescan:
>>> L = lum.signals.LumiSpectrum(np.arange(25).reshape((5,5)))
>>> lum.savetxt(L, 'linescan.txt')
0.00000  0.00000  1.00000  2.00000  3.00000  4.00000
0.00000  0.00000  5.00000 10.00000 15.00000 20.00000
2.00000  2.00000  7.00000 12.00000 17.00000 22.00000
3.00000  3.00000  8.00000 13.00000 18.00000 23.00000
```

`lumispy.utils.io.to_array(S, axes=True, transpose=False)`

Returns signal object as numpy array (optionally including the axes).

Returns single spectra as two-column array. Returns linescan data as array with signal axis as first row and navigation axis as first column (flipped if `transpose=True`). Returns image as array with the navigation axes as first column and first row. Returns 2D data (e.g. map of a fit parameter value) as array with the signal axes as first column and first row.

**Parameters**

- `axes` *(bool, optional)* – If True (default), include axes in array. If False, return the data array only.
- `transpose` *(bool, optional)* – If True, transpose data array and exchange axes. Default is false. Ignored for single spectra.
- `**kwargs` – Takes any additional arguments of `numpy.loadtxt`, e.g. `newline header, footer, comments, or encoding`.

**Notes**

The output of this function can be used to convert a signal object to a pandas dataframe, e.g. using `df = pd.DataFrame(lum.to_array(S))`. 
Examples

```python
>>> import lumispy as lum
>>> import numpy as np
...
>>> # Spectrum:
>>> S = lum.signals.LumiSpectrum(np.arange(5))
>>> lum.to_array(S)
array([[0., 0.],
       [1., 1.],
       [2., 2.],
       [3., 3.],
       [4., 4.]])
...
>>> # Linescan:
>>> L = lum.signals.LumiSpectrum(np.arange(25).reshape((5,5)))
>>> lum.to_array(L)
array([[ 0., 0., 1., 2., 3., 4.],
       [ 0., 0., 1., 2., 3., 4.],
       [ 1., 5., 6., 7., 8., 9.],
       [ 2., 10., 11., 12., 13., 14.],
       [ 3., 15., 16., 17., 18., 19.],
       [ 4., 20., 21., 22., 23., 24.]])
```

lumispy.utils.signals module

lumispy.utils.signals.com(spectrum_intensities, signal_axis, **kwargs)

Finds the centroid (center of mass) of a peak in the spectrum based from the intensity at each pixel value and its respective signal axis.

Parameters

- **spectrum_intensities** (array) – An array with the intensities of the spectrum.
- **signal_axis** (hyperspy.axes.BaseDataAxis subclass) – A HyperSpy signal axis class containing an array with the wavelength/energy for each intensity/signal value.
- **kwargs** (dictionary) – For the scipy.interpolate.interp1d function.

Returns

- **center_of_mass** – The centroid of the spectrum.

Return type

float
Examples

```python
# Assume we have a spectrum with wavelengths and intensities >>> wavelengths = [200, 300, 400, 500, 600, 700] >>> intensities = [1, 2, 3, 2, 1, 0] >>> from hyperspy.axes import DataAxis >>> signal_axis = DataAxis(axis=wavelengths)

>>> center_of_mass = com(intensities, signal_axis)
>>> print(center_of_mass)  # Outputs: [400.0]
```

Module contents

1.10 Changelog

All notable changes to this project will be documented in this file.
The format is based on Keep a Changelog, and this project adheres to Semantic Versioning.

1.10.1 2023-03-15 - version 0.2.2

Changed

- Use GitHub code scanning (CodeQL) for integrity check as it replaces LGTM
- Added a centroid/center of mass functionality to analyse peak position of a spectrum (both in `utils` and in `LumiSpectrum`)
- Add documentation of signal tools

Maintenance

- Replace `sphinx.ext.imgmath` by `sphinx.ext.mathjax` to fix the math rendering in the `ReadTheDocs` build
- fix external references in the documentation

1.10.2 2022-11-02 - version 0.2.1

Added

- Improved documentation
- Use lgtm.com to check code integrity
Changed

- Fix conversion to Raman shift (relative wavenumber) and make jacobian=False default; fix inplace=False for axis conversions
- Fix to_eV and to_invcm, as slicing with .isig[ ] was failing on converted signals
- s.remove_negative now defaults to inplace=False (previously True)

Maintenance

- Use softprops/action-gh-release action instead of deprecated create-release, pin action to a commit SHA

1.10.3 2022-04-29 - version 0.2.0

Added

- Set up read the docs documentation
- Added metadata convention
- Add proper handling of variance on Jacobian transformation during axis conversion (eV, invcm)

Changed

- Account for the general availability of non-uniform axes with the HyperSpy v1.7 release
- Make LumiTransient 1D and add 2D LumiTransientSpectrum class
- Add python 3.10 build, remove python 3.6
- Fix error in background dimensions that allows compatibility for updated map in HyperSpy (failing integration tests)
- Fix for links in PyPi
- Deprecate exposure argument of s.scale_by_exposure in favor of integration_time in line with metadata convention
- Add deprecation warning to remove_background_from_file

1.10.4 2021-11-23 - version 0.1.3

Changed

- Mentions of the now deleted non_uniform_axes branch in HyperSpy updated to RELEASE_next_minor
- Change ‘master’ to ‘main’ branch
- Updated/corrected badges and other things in README.md and other documentation files
1.10.5 2021-08-22 - version 0.1.2

Added

• This project now keeps a Changelog
• Added signal-hierarchy for time-resolved luminescence
• Added GitHub action for release
• Created logo

Changed

• Consistent black-formatting
• fixed join_spectra
• fixed tests

1.10.6 2021-03-26 - version 0.1.0

Added

• The first release, basic functionality implemented

1.11 Citing LumiSpy

LumiSpy is maintained by an active community of developers.

If LumiSpy has been significant to a project that leads to an academic publication, please acknowledge that fact by citing it. The DOI in the badge below is the Concept DOI – it can be used to cite the project without referring to a specific version. If you are citing LumiSpy because you have used it to process data, please use the DOI of the specific version that you have employed. You can find it by clicking on the DOI badge below:

1.12 Contributing

LumiSpy is meant to be a community maintained project. We welcome contributions in the form of bug reports, documentation, code, feature requests, and more. In the following we refer to some resources to help you make useful contributions.
1.12.1 Issues

The issue tracker can be used to report bugs or propose new features. When reporting a bug, the following is useful:

- give a minimal example demonstrating the bug,
- copy and paste the error traceback.

1.12.2 Pull Requests

If you want to contribute to the LumiSpy source code, you can send us a pull request. Small bug fixes are corrections to the user guide are typically a good starting point. But don’t hesitate also for significant code contributions - if needed, we’ll help you to get the code ready to common standards.

Please refer to the HyperSpy developer guide in order to get started and for detailed contributing guidelines.

The kikuchipy contributors guide, another HyperSpy extension, also is a valuable resource that can get you started and provides useful guidelines.

Reviewing

As quality assurance, to improve the code, and to ensure a generalized functionality, pull requests need to be thoroughly reviewed by at least one other member of the development team before being merged.

1.12.3 Documentation

The LumiSpy documentation consists of three elements:

- Docstrings following the numpy standard that document the functionality of individual methods on GitHub.
- The documentation written using Sphinx and hosted on Read the Docs. The source is part of the GitHub repository.
- A set of curated Jupyter notebooks in the LumiSpy demos repository on GitHub that provide tutorials and example workflows.

Improving documentation is always welcome and a good way of starting out to learn the GitHub functionality. You can contribute through pull requests to the respective repositories.

1.12.4 Code style

LumiSpy follows Style Guide for Python Code with The Black Code style.

For docstrings, we follow the numpydoc standard.

Package imports should be structured into three blocks with blank lines between them:

- standard libraries (like os and typing),
- third party packages (like numpy and hyperspy),
- and finally lumispy imports.
1.12.5 Writing tests

All functionality in LumiSpy is tested via the pytest framework. The tests reside in the test directory. Tests are short methods that call functions in LumiSpy and compare resulting output values with known answers. Please refer to the HyperSpy development guide for further information on tests.

1.12.6 Releasing a new version

LumiSpy versioning follows semantic versioning and the version number is therefore a three-part number: MAJOR.MINOR.PATCH. Each number will change depending on the type of changes according to the following:

- MAJOR increases when making incompatible API changes,
- MINOR increases when adding functionality in a backwards compatible manner, and
- PATCH increases when making backwards compatible bug fixes.

The process to release a new version that is pushed to PyPI and Conda-Forge is documented in the Releasing guide.

1.13 License

LumiSpy is free software: you can redistribute it and/or modify it under the terms of the GNU General Public License (GPL) as published by the Free Software Foundation, either version 3 of the license, or (at your option) any later version. LumiSpy is distributed in the hope that it will be useful, but \textbf{without any warranty}; without even the implied warranty of \textbf{merchantability} or \textbf{fitness for a particular purpose}. See the GNU General Public License for more details.


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