

Indirect Hard Modelling, in Python

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# Contents

1	Int	roduction
2	Us 2.1 2.2 2.3 2.4	er guide Installation
3	Lis	st of modules and functions
•	3.1	MODULE input reading
	0.1	3.1.1 input_reading.read_input
		3.1.2 input reading.read input file
	3.2	MODULE spectra_reading
		3.2.1 spectra reading.Multiplet
		3.2.2 spectra reading.Spectr
		3.2.3 spectra_reading.main
	3.3	MODULE gen_param
		3.3.1 gen_param.L2P
		3.3.2 gen_param.P2L
		3.3.3 gen_param.as_par
		3.3.4 gen_param.main
		$3.3.5$ gen_param.multiplet2par
		3.3.6 gen_param.singlet2par
	3.4	MODULE fit_mixture
		3.4.1 fit_mixture.L2P
		3.4.2 fit_mixture.calc_spectra
		3.4.3 fit_mixture.calc_spectra_obj
		3.4.4 fit_mixture.f2min
		3.4.5 fit_mixture.main
		3.4.6 fit_mixture.main
		3.4.7 fit_mixture.write_output
	3.5	MODULE plots
		3.5.1 plots.convergence_path
		3.5.2 plots.plot_iguess
		3.5.3 plots.plot_output

# 1. Introduction

pyIHM is a python software designed in order to offer a comprehensive interface to perform quantitative analyses on NMR spectra of mixtures, using the Indirect Hard Modelling<sup>1</sup> approach.

The Indirect Hard Modelling consists into performing a deconvolution of the spectrum of the mixture using the spectra of the individual components as basis set. Conceptually, the algorithm is made of four steps:

- 1. fit the spectra of the components of the mixture with a hard model (e.g. Voigt);
- 2. read and process the spectrum of the mixture;
- 3. make the initial guess using the set of peaks generated at point 1;
- 4. get the relative concentrations of the components in the mixture.

The routines for reading and processing of the spectra and for the generation of the models rely on the KLASSEZ<sup>2</sup> package.

<sup>&</sup>lt;sup>1</sup>Ernesto Kriesten et al. "Fully automated indirect hard modeling of mixture spectra". In: *Chemometrics and Intelligent Laboratory Systems* 91.2 (2008), pp. 181–193; Anton Duchowny et al. "Quantification of PVC plasticizer mixtures by compact proton NMR spectroscopy and indirect hard modeling". In: *Analytica Chimica Acta* 1229 (2022), p. 340384.

 $<sup>^2\</sup>mathit{KLASSEZ: a\ package\ for\ the\ management\ of\ NMR\ data.\ 2023.\ \mathtt{URL:\ https://github.com/MetallerTM/klassez.}}$ 

# 2. User guide

#### 2.1 Installation

pyIHM can be installed from the associated PyPI repository using pip from the command line by typing:

```
pip install pyihm
```

Alternatively, it is possible to download the .whl file, located in the dist/ folder of the GitHub repository, and install it with pip:

pip install <filename>.whl

## 2.2 Write the input file

Once installed, the software can be operated from the command line via typing:

```
python3 -m pyihm <input_file>
```

where <input\_file> is the path to the input file that contains the parameters for the run. Multiple input files can be given at once, writing their paths in sequence without punctuation signs between them.

```
python3 -m pyihm <input_file_1> <input_file_2> <input_file_3>
```

The input file must be written in a plain text file. It consists in a series of keywords, followed by their arguments in the following line. The sections of the file, one per keyword, are separated by an empty line.

A template for the input file is shown in table 2.1. A detailed explanation of the keyword meanings and the syntax of the related parameters follows.

#### • BASE\_FILENAME

Root of the name of all files that the program will save.

#### • MIX\_PATH

Path to the input spectrum (raw). The folder/file to be read is the first argument, followed by comma-separated additional parameters. It is very important to specify the spectrometer format to allow proper reading, using the <code>spect='format'</code> keyword. The accepted formats are: <code>bruker</code> for Bruker, <code>varian</code> for <code>Varian/Agilent</code>, <code>magritek</code> for SpinSolve benchtop, <code>oxford</code> for Oxford Instruments and general .jdx files.

#### • MIX\_SPECTRUM\_TXT—optional

Path to a plain text file that contains the intensity values of the real part of the spectrum. Useful to be set if the mixture spectrum was processed with anexternal software.

Table 2.1: Example of the input file, used to run the test located in the test/ folder of the GitHub repository.

```
BASE_FILENAME
output/test
MIX_PATH
M.acqus, isexp=False
MIX_SPECTRUM_TXT
M.r
COMP_PATH
comp/C_1.fvf
comp/C_2.fvf
comp/C_3.fvf
FIT_LIMITS
10, 0
FIT_BDS
utol=0.2
utol_sg=0.01
stol=0.01
ktol=0.001
PLT_OPTS
ext=tiff, dpi=300
```

#### • COMP\_PATH

List of the .fvf files that contain the parameters of the signals of the components, generated by KLASSEZ. See section 2.3 for details. Write one file per row.

#### • FIT\_LIMITS

Limits of the fitting region, in ppm, separated by a comma.

#### • FIT\_BDS

Tolerances for the parameters during the fit.

- utol: tolerance for the chemical shifts, in ppm. Given the starting chemical shift  $\delta$ , they will vary in the interval  $[\delta \text{utol}, \delta + \text{utol}]$ .
- utol\_sg: tolerance for the chemical shifts of the signals within the same group, in ppm. Given the starting chemical shift  $\delta$ , they will vary in the interval  $[\delta \text{utol\_sg}, \delta + \text{utol\_sg}]$ .
- stol: tolerance for the linewidths. They will vary in the interval  $[\delta-\mathtt{utol}, \delta+\mathtt{utol}]$ . Given the starting linewidth  $\Gamma$ , in Hz, they will vary in the interval  $[\Gamma-\mathtt{stol}\cdot\Gamma,\Gamma+\mathtt{stol}\cdot\Gamma]$ .
- ktol: tolerance for the relative intensities of the signals in the same spectrum. Given the starting intensity k, they will vary in the interval [k ktol, k + ktol].

#### • FIT\_KWS—optional

Additional parameters for lmfit.Minimizer.minimize, except the 'method' key, which is always set as 'nelder'.

• PLT\_OPTS—optional
Set specific resolution (dpi) and format (ext) for the figures that will be saved. The default values are ext='tiff', dpi=600.

## 2.3 Deconvolution of the spectral components

A script for the deconvolution of the spectra is provided here. It uses the KLASSEZ package to read, process and deconvolve the spectra. The script must be edited in order to fit the specific user's need.

```
#! /usr/bin/env python3
import sys
import klassez as kz
filename = sys.argv[1] # Spectrum
spect = sys.argv[2]
                      # Format
# Read the spectrum
S = kz.Spectrum_1D(filename, spect=spect)
# Do FT
S.process()
# Set "if 1" to phase correct
if 0:
   S.adjph()
# Create/read the initial guess for the deconvolution
S.F.iguess()
# Perform the fit...
S.F.dofit( # ...with the following options:
       u_tol=0.2,
                         # variation on chemical shift /ppm
       f_tol=2,
                          # variation of FWHM /Hz
       vary_phase=False, # Phase correction on the peaks
       vary_xg=False,
                          # Fraction of gaussianity
# Save the figures for the fit
S.F.plot('result', show_res=True, res_offset=0.1)
```

At the end of the run, the .fvf file will be saved in the folder of the spectrum.

# 2.4 Reading the results

During the execution of pyIHM, a series of files will be generated. The .out file contains a summary of the result of the fit, i.e. the concentrations of the species of the mixture and the parameters of the individual signals.

Regarding the figures, the '\_iguess' one diplays the initial guess for the fit, whereas the '\_total' one shows the result. The '\_wcomp' figure highlights the components with different colors. Finally, the '\_rhist' figure is the histogram of the residuals, with a gaussian curve overlayed in red. The statistical parameters of the residuals (i.e. the mean and the standard deviation) are reported in the legend of this figure.

# 3. List of modules and functions

# 3.1 MODULE input reading

## 3.1.1 input reading.read input(filename)

Reads the input file to get all the information to perform the fit. The values read from the file are double-checked, and the missing entries are replaced with default values, so not to leave space to stupid mistakes.

#### **Parameters:**

• filename: strPath to the input file

#### Returns:

- base\_filename: strRoot of the name of all the files that the program will save
- mix\_path: str
  Path to the mixture spectrum
- mix\_kws: dict of keyworded arguments
  Additional instructions to be passed to kz.Spectrum\_1D.\_\_init\_\_
- mix\_spectrum\_txt: str or None
  Path to a .txt file that contains a replacement spectrum for the mixture
- comp\_path: *list*Path to the .fvf files to be used for building the spectra of the components
- fit\_lims: tuple
  Limits of the fitting region, in ppm
- fit\_bds: dict
  Boundaries for the fitting parameters. The keywords are:
  - utol = allowed displacement for singlets and whole multiplets, in ppm (absolute)
  - utol\_sg = allowed displacement for the peaks that are part of the same multiplet relatively to the center, in ppm (absolute)
  - stol = allowed variation for the linewidth, in Hz (relative)
  - ktol = allowed variation for the relative intensities within the same spectrum(relative)

#### 6

# 3.1.2 input\_reading.read\_input\_file(filename)

Runs over the input file, looks for specific keywords, and interpret them accordingly.

#### Parameters:

• filename: strPath to the input file

#### Returns:

• dic: dict Read values, organized

#### 7

## 3.2 MODULE spectra reading

### 3.2.1 spectra reading.Multiplet

class

Class that represent a multiplet as a collection of peaks.

#### Attributes:

- acqus: dict
  Dictionary of acquisition parameters
- peaks: *dict*Dictionary of kz.fit.Peak objects
- U: float
  Mean chemical shift of the multiplet
- u\_off: dict Chemical shift of the components of the multiplet, expressed as offset from self.U

#### Methods:

\_\_init\_\_(self, acqus, \*peaks)

Initialize the class.

#### Parameters:

- acqus: *dict*Dictionary of acquisition parameters
- peaks: kz.fit.Peak objects

  Peaks that are part of the multiplet. They must have an attribute 'idx' which serves as label

 $\_\_call\_\_(self)$ 

Compute the trace correspondant to the multiplet.

#### Returns:

• trace: *1darray* Sum of the components

#### par(self)

Computes a summary dictioanary of all the parameters of the multiplet.

#### **Returns:**

• dic: dict of dict

The keys of the inner dictionary are the parameters of each single peak, the outer keys are the labels of the single components

### 3.2.2 spectra reading.Spectr

class

Class that represents a spectrum as a collection of peaks and multiplets.

#### Attributes:

- acqus: *dict*Acquisition parameters
- peaks: dict
  Dictionary of peaks object, labelled according to the 'idx' attribute of each single peak
- unique\_groups: *list*Identifier labels for the multiplets, without duplicates
- p\_collections: dict
  Dictionary of kz.fit.Peak and Multiplet objects, labelled according to the group they belong
  to. In particular, self.p\_collections[0] is a list of kz.fit.Peak objects, whereas all the remaining
  entries consist of a single Multiplet object.
- total: *1darray*Placeholder for the trace of the spectrum, as sum of all the peaks.

#### Methods:

\_\_init\_\_(self, acqus, \*peaks)

Initialize the class.

#### Parameters:

- acqus: dict
  Dictionary of acquisition parameters
- peaks: kz.fit.Peak objects

  Peaks that are part of the multiplet. They must have an attribute 'idx' which serves as label

$$\_\_call\_\_(self, I=1)$$

Compute the total spectrum, multiplied by I.

#### Parameters:

• I: *float*Intensity value that multiplies the spectrum

#### **Returns:**

• total: *1darray*Computed spectrum

#### calc total(self)

Computes the sum of all the peaks to make the spectrum

## Returns:

• total: *1darray* Computed spectrum

### 3.2.3 spectra reading.main(M, spectra dir, lims=None)

Reads the .fvf files, containing the fitted parameters of the peaks of a series of spectra. Then, computes a list of Spectr objects with those parameters, and returns it. The relative intensities are referred to the total intensity of the whole spectrum, not to the ones of the fitted regions. Employs kz.fit.read\_vf to read the .fvf files and generate the parameters.

#### Parameters:

- M: kz.Spectrum\_1D object Mixture spectrum. Used to get the spectral parameters for the kz.fit.Peak objects
- spectra\_dir: *list of str*Sequence of the locations of the .fvf files to be read
- lims: *tuple*Borders of the fitting window, in ppm (left, right)

#### Returns:

• collections: *list of Spectr objects*Spectra of pure components, treated as collections of peaks.

# 3.3 MODULE gen param

## 3.3.1 gen param.L2P(L, Xmin, Xmax)

Convert a normalized parameter into its absolute counterpart.

#### Parameters:

• L: *float*Normalized parameter value

• Xmin: *float*Lower bound of the 'original' patameter

• Xmax: *float*Upper bound of the 'original' patameter

#### Returns:

• name: str Label of the parameter

• value: float

Correspondant value

## 3.3.2 gen param.P2L(P)

Normalize a lmfit. Parameter object according to its boundaries. Works only if expr is not set! In this case, in fact, it returns None. The boundaries of the new parameter are set to be (0,1), where 0 corresponds to P.min and 1 to P.max.

#### Parameters:

• P: *lmfit.Parameter object* Not normalized parameter

#### Returns:

• L: *lmfit.Parameter object* Normalized parameter. If P.expr is set, this is None.

### 3.3.3 gen param.as par(name, value, lims=0, rel=True)

Creates a lmfit. Parameter object using the given parameters.

#### Parameters:

- name: strLabel of the parameter
- value: float or str

  If it is float, it is the value of the parameter. If it is a str, it is put in the 'expr' attribute of the lmfit. Parameter object.
- lims: float or tuple
  Determines the boundaries. If it is a tuple, the boundaries are min(lims) and max(lims). If it is a single float, the boundaries are (value-lims, value+lims). Not read if value is str
- rel: bool Relative boundaries. If it is True and lims is a float, the boundaries are set to value-lims\*value, value+lims\*value.

#### **Returns:**

• p: lmfit.Parameter object
Object created according to the given parameter

## 3.3.4 gen param.main(M, components, bds)

Create the lmfit.Parameters objects needed for the fitting procedure.

#### Parameters:

- M: kz.Spectrum\_1D object Mixture spectrum
- components: *list*List of Spectra objects
- bds: *dict*Boundaries for the fitting parameters.

#### Returns:

- Lparam: *lmfit.Parameters object* Normalized parameters for the fit
- param: *lmfit.Parameters object* Actual parameters for the fit

### 3.3.5 gen param.multiplet2par(item, spect, group, bds)

Converts a Multiplet object into a list of lmfit.Parameter objects. The keys are of the form 'S $\#_p$ ?' where # is spect and ? is the index of the peak. p = U is the mean chemical shift p = o is the offset from U p = u is the absolute chemical shift, computed as U + o, set as expression.

#### Parameters:

• item: fit.Peak object
Peak to convert into Parameter. Make sure the .idx attribute is set!

• spect: *int*Label of the spectrum to which the peak belongs to

• group: *int*Label of the multiplet group

• bds: *dict*Contains the parameters' boundaries

#### Returns:

• p: *list*List of lmfit.Parameter objects

## 3.3.6 gen param.singlet2par(item, spect, bds)

Converts a fit.Peak object into a list of lmfit.Parameter objects: the chemical shift (u), the linewidth (s), and intensity (k). The keys are of the form 'S#\_p?' where # is spect and ? is the index of the peak.

#### Parameters:

• item: kz.fit.Peak object Peak to convert into Parameter. Make sure the .idx attribute is set!

• spect: *int*Label of the spectrum to which the peak belongs to

• bds: *dict*Contains the parameters' boundaries

#### Returns:

• p: *list*List of lmfit.Parameter objects

# 3.4 MODULE fit\_mixture

## 3.4.1 fit mixture.L2P(L, Xmin, Xmax)

Convert a normalized parameter into its absolute counterpart.

#### Parameters:

• L: *float*Normalized parameter value

• Xmin: *float*Lower bound of the 'original' patameter

• Xmax: *float* Upper bound of the 'original' patameter

#### Returns:

• name: str Label of the parameter

• value: float

Correspondant value

# $3.4.2 \quad fit\_mixture.calc\_spectra(Lparam, \, param, \, N\_spectra, \, acqus, \, N)$

Computes the spectra to be used as components for the fitting procedure, in form of lists of 1darrays. Each array is the sum of all the peaks. This function is called at each iteration of the fit.

#### Parameters:

• Lparam: *lmfit.Parameters object* Normalized parameters

• param: *lmfit.Parameters object* Actual parameters

• N\_spectra: *int* Number of spectra to be used as components

• acqus: *dict*Dictionary of acquisition parameters

• N: *int*Number of points for zero-filling, i.e. final dimension of the arrays

#### Returns:

• spectra: *list of 1darray*Computed components of the mixture, weighted for their relative intensity

# 3.4.3 fit\_mixture.calc\_spectra\_obj(Lparam, param, N\_spectra, acqus, N)

Computes the spectra to be used as components for the fitting procedure, in form of lists of kz.fit.Peak objects.

#### Parameters:

- Lparam: *lmfit.Parameters object* Normalized parameters
- param: *lmfit.Parameters object* Actual parameters
- N\_spectra: *int* Number of spectra to be used as components
- acqus: *dict*Dictionary of acquisition parameters
- N: *int*Number of points for zero-filling, i.e. final dimension of the arrays

#### **Returns:**

• spectra: *list of kz.fit.Peak objects*Computed components of the mixture, weighted for their relative intensity

# 3.4.4 fit\_mixture.f2min(Lparam, param, N\_spectra, acqus, N, exp, I, plims)

Function to compute the quantity to be minimized by the fit.

#### Parameters:

- Lparam: *lmfit.Parameters object* Normalized parameters
- param: *lmfit.Parameters object* actual parameters
- N\_spectra: *int*Number of spectra to be used as components
- acqus: *dict*Dictionary of acquisition parameters
- N: *int*Number of points for zero-filling, i.e. final dimension of the arrays
- exp: *1darray* Experimental spectrum
- I: float
  Intensity correction for the calculated spectrum. Used to maintain the relative intensity small.
- plims: *slice*Delimiters for the fitting region. The residuals are computed only in this regio. They must be given as point indices

#### Returns:

• target: float
Given the experimental data y and the calculated data  $y_c$ :  $\sum [(y - I * y_c)^2]$ 

## 3.4.5 fit mixture.main(M, components, bds)

Create the lmfit.Parameters objects needed for the fitting procedure.

#### Parameters:

- M: kz.Spectrum\_1D object Mixture spectrum
- components: *list*List of Spectra objects
- bds: *dict*Boundaries for the fitting parameters.

#### Returns:

- Lparam: *lmfit.Parameters object* Normalized parameters for the fit
- param: *lmfit.Parameters object* Actual parameters for the fit

# 3.4.6 fit\_mixture.main(M, N\_spectra, Lparam, param, lims=None, file-name='fit', ext='tiff', dpi=600)

Core of the fitting procedure. It computes the initial guess, save the figure, then starts the fit. After the fit, writes the output file and saves the figures of the result. Summary of saved files:

- '<filename>.out': fit report
- '<filename> iguess.<ext>': figure of the initial guess
- '<filename>\_total.<ext>': figure that contains the experimental spectrum, the total fitting function, and the residuals
- '<filename>\_wcomp.<ext>': figure that contains the experimental spectrum, the total fitting function, and the components in different colors. The residuals are not shown
- '<filename>\_rhist.<ext>': histogram of the residual, with a gaussian function drawn on top according to its statistical parameters.

#### Parameters:

- M: kz.Spectrum\_1D object Mixture spectrum
- N\_spectra: *int* Number of spectra to be used as fitting components
- Lparam: *lmfit.Parameters object* Normalized parameters
- param: *lmfit.Parameters object* Actual parameters
- lims: tuple or None
  Delimiters of the fitting region, in ppm. If None, the whole spectrum is used.
- filename: strRoot of the names for the names of the files that will be saved.
- ext: strFormat of the figures
- dpi: *int*Resolution of the figures, in dots per inches

## 3.4.7 fit mixture.write output(M, I, K, spectra, lims, filename='fit.report')

Write a report of the performed fit in a file. The parameters of the single peaks are saved using the kz.fit.write\_vf function.

#### Parameters:

- M: kz.Spectrum\_1D object Mixture spectrum
- I: *float*Absolute intensity for the calculated spectrum
- K: sequence Relative intensities of the spectra in the mixture
- lims: *tuple*Boundaries of the fit region
- filename: strName of the file where to write the files.

# 3.5 MODULE plots

# 3.5.1 plots.convergence\_path(conv\_path, filename='conv', ext='tiff', dpi=600)

Makes the figures of the final fitted spectrum and saves them. Three figures are made: look at the fitting.main function documentation for details.

#### Parameters:

• conv\_path: strPath to the file of the convergence path

• filename: strFilename of the final figure

• ext: strFormat of the figure

• dpi: *int*Resolution of the figure, in dots per inches

# 3.5.2 plots.plot\_iguess(ppm\_scale, exp, total, components, lims=None, X label='\$\delta\$ /ppm', filename='fit', ext='tiff', dpi=600)

Makes the figure of the initial guess and saves it.

#### Parameters:

- ppm\_scale: 1darray
  PPM scale of the spectrum
- exp: *1darray*Mixture spectrum, real part
- total: *1darray* Fitting function
- components: *list of 1darray*Spectra used as components, real part
- lims: tuple or None
  Delimiters of the fitting region, in ppm. If None, the whole spectrum is used.
- X\_label: strLabel for the X\_axis
- filename: strThe name of the figure will be <filename> iguess.<ext>
- ullet ext: str Format of the figures
- dpi: *int*Resolution of the figures, in dots per inches

# 3.5.3 plots.plot\_output(ppm\_scale, exp, total, components, lims=None, X label=' $\frac{ppm'}{lense}$ , filename='fit', ext='tiff', dpi=600)

Makes the figures of the final fitted spectrum and saves them. Three figures are made: look at the fitting.main function documentation for details.

#### Parameters:

- ppm\_scale: 1darray
  PPM scale of the spectrum
- exp: *1darray*Mixture spectrum, real part
- total: *1darray*Fitting function
- components: *list of 1darray*Spectra used as components, real part
- lims: tuple or None
  Delimiters of the fitting region, in ppm. If None, the whole spectrum is used.
- X\_label: strLabel for the X\_axis
- filename: strRoot filename for the figures
- ext: strFormat of the figures
- dpi: *int*Resolution of the figures, in dots per inches