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# 1. Introduction

KLASSEZ is a python package written to handle 1D and 2D NMR data. The aim of the project is to provide a toolkit, consisting of 'black-box' functions organized in modules, that could be used to read, process and analyze such data in a flexible manner, so to adapt to the needs of the individual users. However, the open-source nature of the package grants the user the chance to open the lid of these black-boxes and understand the gears that stand behind the function call.

The development of the toolkit started with python 3.8 and therefore it is compatible with that version. Nevertheless, the use of python 3.10 is advised.

The key objects provided by KLASSEZ are the classes Spectrum\_1D and Spectrum\_2D, that are able to fulfil the aims of the package with a few lines of code. The classes are able to read both simulated (i.e. generated with a custom-made input file) and experimental datasets. The latter feature was tested with Bruker data after the removal of the digital filter (run command convdta in TopSpin), but should be compatible with other kind of spectrometers, thanks to the remarkable work made by J. J. Helmus and coworkers with their nmrglue package<sup>1</sup>. Either the FID or the spectrum processed with external solver can be read from KLASSEZ by using the classes Spectrum\_nD or pSpectrum\_nD, respectively.

The processing module, besides the classical functions used for the processing of NMR data (window functions, Fourier transform, etc.), includes denoising algorithms based on Multivariate Curve Resolution<sup>2</sup> and on Cadzow method<sup>3</sup>. Details are illustrated in the description of the functions.

Functions to show and analyze data in real time are provided, with dedicated GUIs. However, it is better to rely on the standalone functions, enclosed in the single modules, to save the figures. In fact, the figures module offers a wide plethora of functions (all based on matplotlib) to plot the data with a high degree of customization for the appearance.

The fitting functions use lmfit to build the initial guess and to minimize the difference between the experimental data and the model, generated with a Voigt profile in the time domain and then Fourier-transformed, in the least-square sense (employing the Levenberg-Marquardt algorithm implemented in scipy). For this purpose, the class Voigt\_fit of the fit module includes attribute functions to construct an initial guess interactively, fit the data, and save the parameters in dedicated files.

Regarding the development of the package, I would like to acknowledge Letizia Fiorucci for her contribution in the design and the implementation of several functions, and for the alpha-testing.

<sup>1</sup>https://www.nmrglue.com/

<sup>&</sup>lt;sup>2</sup>Multivariate Curve Resolution: 50 years addressing the mixture analysis problem - A review

<sup>&</sup>lt;sup>3</sup>Denoising NMR time-domain signal by singular-value decomposition accelerated by graphics processing units

# 2. User guide

### 2.1 Initialize the package

KLASSEZ can be installed from PyPI through:

```
pip install klassez
```

The required dependencies are sorted out automatically.

Initialize the package by writing, at the top of your file:

```
from klassez import *
```

This line executes the following code:

```
import os
import sys
import numpy as np
from numpy import linalg
from scipy import stats
import matplotlib
import matplotlib.pyplot as plt
import matplotlib.cm as cm
import seaborn as sns
from copy import deepcopy
from pprint import pprint

from . import fit, misc, sim, figures, processing
from .Spectra import Spectrum_1D, pSpectrum_1D, Spectrum_2D, pSpectrum_2D
# Use seaborn's colormaps and save it to a dictionary
from .config import CM, CM_2D, COLORS, cron
```

This means these can be not imported in your code, as KLASSEZ already does it for you. An alternative, safer version to prevent overwriting of custom functions is:

```
import klassez as kz
```

In this case, additional packages for the main script must be declared explicitely.

#### 2.1.1 Extra variables

Initializing KLASSEZ also grants access to CM and COLORS.

CM is a dictionary of colormaps taken from seaborn and saved in a dictionary whose keys are their names, so that also matplotlib can use them. You can inspect the keys through:

```
print(CM.keys())
```

There is a restricted list of color maps, CM\_2D, that should be used for visualizing 2D spectra. COLORS is:

```
colors = [ 'tab:blue', 'tab:red', 'tab:green', 'tab:orange', 'tab:cyan', 'tab:purple',
    'tab:pink', 'tab:gray', 'tab:brown', 'tab:olive', 'salmon', 'indigo', 'm', 'c', 'g',
    'r', 'b', 'k', ]
```

repeated cyclically ten times and stored as tuple.

Other two 'quality of life' variables are figures.figsize\_small and figures.figsize\_large, which correspond to figure panel sizes of  $3.59 \times 2.56$  inches and  $15 \times 8$  inches, respectively. The former suits well for saving figures of spectra with font sizes of about 10 pt, whereas the latter are best for GUIs and withstand font sizes of about 14 pt.

For NMR: the variable sim.gamma is a dictionary containing the gyromagnetic ratio, in MHz/T, of all the magnetically-active nuclei. For instance:

```
print(sim.gamma['13C'])
>>> 10.70611
```

A decorator function called **cron** is defined in the top-level script **config**, and imported by **\_\_init\_\_**, so that you can use it after writing:

```
from klassez import cron
```

This decorator allows to measure the runtime of a function, and print it on standard output once it ended.

### 2.2 Processing of a 'raw' 1D spectrum

Let us say that your spectrum is saved in the folder /home/myself/spectra/mydataset/1/. Initialize the spectrum object through:

```
Path = "/home/myself/spectra/mydataset/1/"
s = Spectrum_1D(Path)
```

This command will do three main tasks:

- read the binary FID of your spectrum and store it in a complex array s.fid;
- load the acquisition parameters, read the interesting keys and store them in a dictionary s.acqus;
- initialize a dictionary s.procs which contains the processing parameters.

KLASSEZ is able to read also Varian and Spinsolve (Magritek) data, by specifying the option 'spect'.

A detailed description of acqus and procs is shown in table 2.1 and table 2.2.

Please note that reading the spectrum causes the program to save a file called 'name.procs', where 'name' is the path name.

To make the Fourier transform of the FID to obtain the spectrum, you must invoke the **process** method, which reads the **procs** dictionary to get the instructions on the processing you want to make on your spectrum. For instance, if you want to obtain a final spectrum of 8k points with an exponential broadening of  $25 \,\mathrm{Hz}$ :

```
s.procs["wf"]["mode"] = "em"
s.procs["wf"]["lb"] = 25
```

```
s.procs["zf"] = 8192
s.process()
s.pknl()  # Tries to remove the digital filter through a first-order phase correction
```

Calling the process method generates new attributes of the class:

- freq: the frequency scale, in Hz;
- ppm: the ppm scale;
- r: the real part of the spectrum;
- i: the imaginary part of the spectrum;
- S: the complex spectrum (S = r + ii).

After the Fourier transform, the process method applies the phase correction and the calibration using the phase angles and the calibration value saved in the *procs* dictionary automatically. This allows the user to not phase their spectra every time, as well as keeping a record of the processing.

If the spectrum requires phase correction, you can perform it interactively:

```
s.adjph()
```

or by passing the phase angles, in degrees, to adjph. Example, if you know you need to phase your spectrum with 30 degrees of  $\phi^{(0)}$  and -55 degrees of  $\phi^{(1)}$  with the pivot set at 7.32 ppm:

```
s.adjph(p0=30, p1=-55, pv=7.32)
```

In both cases, the phase angles are updated in the procs dictionary.

The spectrum can be calibrated using a dedicated GUI:

```
s.cal()
```

or specifying the shift value in ppm or in Hz (in this case, be sure to set the isHz keyword to True).

```
s.cal(-3) # Shift of -3 ppm
s.cal(1000, isHz=True) # Shift of +1 kHz
```

Both ppm and freq are updated according to the given values.

### 2.2.1 The class pSpectrum\_1D

The class Spectrum\_1D does not work if you want to read the processed data directly from TopSpin (or whatever software you used to acquire and process them). Instead, you should use the class pSpectrum\_1D, which is designed to perform exactly this task. It inherits most of the attributes and methods of the Spectrum\_1D class, therefore its usage closely resembles the example reported in the previous section.

Table 2.1: Description of the acqus dictionary of a Spectrum\_1D object.

Key	Explanation		
В0	Magnetic field strength /T		
BYTORDA Endianness of binary data: 0 little endian, 1 big endian			
DTYPA Binary data type: 0 int32, 2 float64			
GRPDLY Number of points of the digital filter			
nuc Observed nucleus			
olp Carrier frequency i.e. center of the spectrum, in ppm			
01	Same as o1p, but in Hz		
$\mathtt{SWp}$	Sweep width, in ppm		
SW	Sweep width, in Hz		
SF01 Larmor frequency of the observed nucleus at field B0			
TD	Number of sampled complex points		
dw	Dwell time, i.e. the sampling interval, in seconds		
AQ Time duration of the FID			
t1	Acquisition timescale		

Table 2.2: Description of the procs dictionary of a Spectrum_1D object.			
Key Explanation			
wf Window function. This is a dictionary itself:			
	• 'mode': choose function between		
	- 'no': no apodization		
	- 'em': exponential		
	- 'sin': sine		
	- 'qsin': squared sine		
- 'gm': mixed lorentzian-gaussian			
- 'gmb': mixed lorentzian-gaussian, Bruker style			
• 'lb': Exponential line-broadening. Read by em and gm			
	• 'lb_gm': Exponential line-broadening. Read by gm		
	• 'gb': Gaussian line-broadening. Read by gmb		
	• 'gb_gm': Gaussian line-broadening. Read by gm		
	• 'gc': Center of the gaussian $\in [0,1]$ . Read by gm		
	• 'ssb': Shift of the sine bell. Read by sin and qsin		
	• 'sw': Sweep width. Automatically set according to acqus['SW']		
zf tdeff fcor p0 p1 pv	Zero-filling. Set the <i>final</i> number of points!  Number of points to be used for processing  Scaling factor for the first point of the FID before Fourier transform  Frequency-independent phase correction /degrees  First order phase correction /degrees  Pivot point for the first order phase correction /ppm		
basl_c			

Offset, in ppm, to be added to the frequency and ppm scales for cali-

the 0-order coefficient  $\,$ 

bration

cal

# 2.3 Processing of a 'raw' 2D spectrum

Let us say that your spectrum is saved in the folder /home/myself/spectra/mydataset/21/. Initialize the spectrum object through:

```
Path = "/home/myself/spectra/mydataset/21/"
s = Spectrum_2D(Path)
```

The generated acqus and procs dictionaries include informations on both dimensions.

Table 2.3: Description of the acqus dictionary of a Spectrum\_2D object.

Key Explanation			
B0	Magnetic field strength /T		
BYTORDA Endianness of binary data: 0 little endian, 1 big endian			
DTYPA	Binary data type: 0 int32, 2 float64		
GRPDLY	Number of points of the digital filter		
nuc1	Observed nucleus in the indirect dimension		
nuc2	Observed nucleus in the direct dimension		
o1p	Carrier frequency i.e. center of the indirect dimension, in ppm		
o2p Carrier frequency i.e. center of the direct dimension, in ppm			
o1 Same as o1p, but in Hz			
o2 Same as o2p, but in Hz			
SW1p Sweep width of the indirect dimension, in ppm			
SW2p Sweep width of the direct dimension, in ppm			
SW1 Sweep width of the indirect dimension, in Hz			
Sweep width of the indirect dimension, in Hz			
SF01 Larmor frequency of the observed nucleus in F1 at field B0			
SF02 Larmor frequency of the observed nucleus in F2 at field B0			
TD1 Number of $t_1$ -increments			
TD2 Number of sampled complex points			
dw1 $t_1$ increments, in seconds			
dw2 Dwell time, i.e. the sampling interval, in seconds			
AQ1 Sampled timescale of the indirect dimension			
AQ2	Time duration of the FID		
t1	Evolution timescale		
t2	Acquisition timescale		

Then, the sequence of commands resembles the ones of the 1D spectra.

The keys for adjph are of the kind: pXY, where X is the order of the phase correction (0 or 1) and Y is the dimension on which to apply it (1 or 2). Explicative table below:

s.process()

s.pknl() # Remove the digital filter

<sup>#</sup> Also in this case, phase correction and calibration are performed automatically with the values in procs

s.adjph()

s.plot()

Table 2.4: Description of the procs dictionary of a Spectrum\_2D object. Each of these dictionary entry is a list of two elements: the first one (index 0) is the processing to apply on the indirect dimension, the second (index 1) on the direct dimension. For instance, procs[tdeff] = [64, 1024] means to truncate the indirect evolutions to 64 points and the FIDs to 1024 points.

Key	Explanation
wf	Window function. This is a dictionary itself:
	• 'mode': choose function between
	- 'no': no apodization
	- 'em': exponential
	- 'sin': sine
	- 'qsin': squared sine
	<ul><li>- 'gm': mixed lorentzian-gaussian</li></ul>
	- 'gmb': mixed lorentzian-gaussian, Bruker style
	• 'lb': Exponential line-broadening. Read by em and gmb
	• 'lb_gm': Exponential line-broadening. Read by gm
	• 'gb': Gaussian line-broadening. Read by gmb
	• 'gb_gm': Gaussian line-broadening. Read by gm
	• 'gc': Center of the gaussian $\in [0,1]$ . Read by gm
	• 'ssb': Shift of the sine bell. Read by sin and qsin
	• 'sw': Sweep width. Automatically set according to acqus['SW']
zf	Zero-filling. Set the <i>final</i> number of points!
tdeff	Number of points to be used for processing
fcor p02	Scaling factor for the first point of the FID before Fourier transform Frequency-independent phase correction /degrees, direct dimension
p12	First order phase correction /degrees, direct dimension
pv2	Pivot point for the first order phase correction /ppm, direct dimensio
p01	Frequency-independent phase correction /degrees, indirect dimension
p11	First order phase correction /degrees, indirect dimension
pv1	Pivot point for the first order phase correction /ppm, indirect dimension
- 4	

Calibration offset for F1 /ppm

Calibration offset for F2 /ppm

cal\_1 cal\_2

	<b>F</b> 1	<b>F2</b>
$\phi^{(0)}$	p01	p02
$\phi^{(1)}$	p11	p12
pivot	pv1	pv2

For further information, rely on the help python builtin function. To read the processed data, use the pSpectrum\_2D class instead.

### 2.3.1 Computing projections

While the 2D spectra give an overall look on the whole experiment, the user might want to extract projection of the direct or the indirect dimension, to focus onto particular features in the spectrum. In order to do so, klassez offers two commands: projf1 and projf2, which compute the sum projections on the indirect or on the direct dimension, respectively, and store the result in dictionaries called trf1 and trf2, whose keys are the ppm values correspondant to the projections. Actually, the capitalized versions of the two dictionaries (with the same keys), i.e. Trf1 and Trf2, can be more useful, as they are instances of the pSpectrum\_1D class and therefore are initialized with ppm scales and other parameters.

Example:

```
# Supposed to have a 1H-15N HSQC spectrum

# Extract the direct dimension trace at 115 ppm, 15N scale
s.projf2(115)

# Access to it through
Proj_115 = s.Trf2['115']

# Extract the indirect dimension trace from 6 to 8 ppm, 1H scale
s.projf1(6, 8)
Proj_indim = s.Trf1['6:8']

# You can plot them:
Proj_115.plot()
Proj_indim.plot()
```

### 2.4 Simulating data

The classes Spectrum\_1D and Spectrum\_2D are also able to generate simulated data by reading a custom-written input file. The functions they use are sim.sim\_1D and sim.sim\_2D.

#### 2.4.1 Simulate 1D data

The input file you have to write *must* have the following keys:

- B0: Magnetic field strength /T;
- nuc: Observed nucleus (e.g. 13C);
- olp: Carrier frequency i.e. centre of the spectrum /ppm;
- SWp: Sweep width /ppm. The spectrum will cover the range [o1p SWp/2, o1p + SWp/2];

- TD: Number of sampled (complex) points;
- shifts: sequence of peak positions /ppm;
- fwhm: Full-width at half-maximum of the peaks /Hz;
- amplitudes: Intensity of the peaks in the FID;
- beta: Fraction of gaussianity.  $\beta=0 \implies$  pure Lorentzian peak,  $\beta=1 \implies$  pure Gaussian peak;

and can have the following keys:

- phases: phases of the peaks /degrees. Default: all zeros;
- mult: fine structures of the peaks (e.g. doublets of triplets: dt). Default: all singlets;
- Jconst: coupling constants of the fine structures /Hz. If more of one coupling is expected, provide them as a sequence. Default: not used as the peaks are all singlets.

Key and value must be separated by a tab character. You are allowed to leave empty rows to improve the readibility and to insert comments using the # character.

Example:

#! /usr/bin/env python3

```
B0 16.4 # 700 MHz 1H
nuc 1H
o1p 4.7
SWp 40
TD 8192

shifts 1, 3, 5, 7
fwhm [10 for k in range(4)]
amplitudes 10, 20, 15, 10
beta 0, 0.4, 0.6, 1
phases 5, 0, 10, 0

mult s, t, dt, ddd
Jconst 0, 15, [12, 9.5], [25, 15, 10]
```

This input file generates the spectrum in Figure 2.1. Code:

```
from klassez import *
s = Spectrum_1D('sim_in_1D', isexp=False)
s.process()
figures.figure1D(s.ppm, s.r, name='test_1D', X_label='$\delta\, ^1$H /ppm',
    Y_label='Intensity /a.u.')
```

### 2.4.2 Simulate 2D data

The same procedure can be followed to simulate 2D spectra. The input file to write is very similar to the one for 1D data, except for the quantities that clearly span over two dimensions. As in NMR textbook, the direct and indirect dimensions will be named F2 and F2 respectively, and dimension-specific quantities will feature the 1 or 2 labels accordingly.

- B0: Magnetic field strength /T;
- nuc1: Observed nucleus in F1(e.g. 13C);
- nuc2: Observed nucleus in F2(e.g. 1H);
- o1p: Carrier frequency i.e. centre of F1 /ppm;
- o2p: Carrier frequency i.e. centre of F2 /ppm;
- SW1p: Sweep width /ppm. The indirect dimension will cover the range [o1p SW1p/2, o1p + SW1p/2];
- SW2p: Sweep width /ppm. The direct dimension will cover the range [o2p SW2p/2, o2p + SW2p/2];
- TD1: Number of sampled (complex) points in F1;
- TD2: Number of sampled (complex) points in F2;
- shifts\_f1: sequence of peak positions in F1 /ppm;
- shifts\_f2: sequence of peak positions in F2 /ppm;
- fwhm\_f1: Full-width at half-maximum of the peaks in F1 /Hz;
- fwhm\_f2: Full-width at half-maximum of the peaks in F2 /Hz;
- amplitudes: Intensity of the peaks in the FID;
- beta: Fraction of gaussianity.  $\beta = 0 \implies$  pure Lorentzian peak,  $\beta = 1 \implies$  pure Gaussian peak;

Phase distortions and fine structures are not allowed for multidimensional spectra. The indirect dimension will be generated employing the *States-TPPI* sampling scheme.

Example:

```
B0 28.2
nuc1
       15N
nuc2
       1H
o1p 115
o2p 5
SW1p
       40
SW2p
       20
TD1 512
TD2 8192
shifts_f1 130.0, 105.0, 120.0, 1.25e2, 130.0, 105.0
shifts_f2 0.0, 0.0, 4.0, 7.0, 1.1e1, 10.5
fwhm_f1 100, 100, 100, 100, 100, 100
fwhm_f2 50, 50, 50, 50, 50, 50
amplitudes 10, 20, 10, 20, 10, 10
       0.0, 0.2, 0.4, 0.6, 0.8, 1.0
beta
```

This input file generates the spectrum in Figure 2.2. Code:

```
#! /usr/bin/env python3
from klassez import *
s = Spectrum_2D('sim_in_2D', isexp=False)
s.process()
figures.figure2D(s.ppm_f2, s.ppm_f1, s.rr, lvl=0.005, name='test_2D', X_label='$\delta\, ^1$H /ppm', Y_label='$\delta\, ^{15}$N /ppm')
```

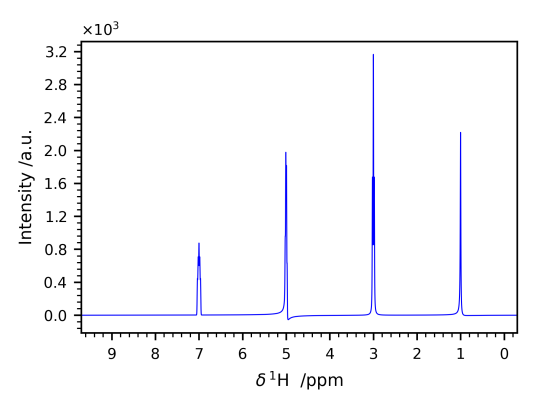


Figure 2.1: Example of a simulated 1D spectrum.

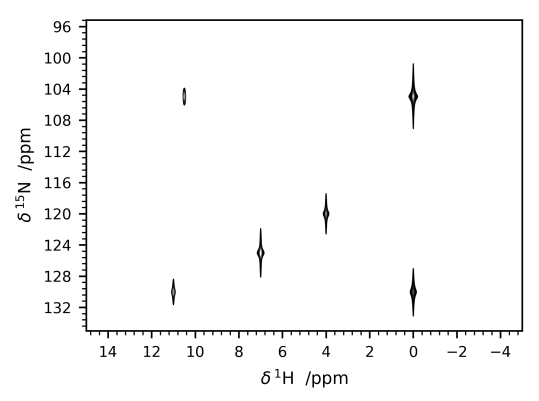


Figure 2.2: Example of a simulated 2D spectrum.

### 2.5 The Pseudo\_2D class

Sometimes, the spectroscopist might find interesting to acquire a series of 1D experiments in which one (or more) parameters are changed according to a certain schedule. This kind of experiments are 2D in principle, but their processing and analysis resemble the one of 1D spectra. Therefore, they lie somewhere in between 1D spectra and 2D spectra, hence they are often referred to as pseudo 2D.

Also in this case, klassez offers a specific class to deal with this kind of data: Pseudo\_2D. Pseudo\_2D is a subclass of Spectrum\_2D; however, many functions have been adapted to resemble the 1D version.

Pseudo\_2D does not encode for a routine to automatically simulate data. If you want to, you should give a 1D-like input file (just like the one in section 2.4.1), and replace the attribute fid with your FID by using the method mount, generated as you wish. With a real dataset this is not required, as it is able to read everything automatically.

```
path_to_pseudo = "/home/myself/spectra/mydataset/899/"
s = Pseudo_2D(path_to_pseudo)
```

The process() function applies apodization, zero-filling and Fourier transform only on the direct dimension, reading the parameters from a procs dictionary like the one of Spectrum\_1D. The attributes freq\_f1 and ppm\_f1 are initialized with np.arange(N), where N is the number of experiments that your FID comprises of. In particular, freq\_f1 numbers the experiments sequentially from 0 to N-1, whereas ppm\_f1 does it from 1 to N. Therefore, when calling the method projf2 to extract the experiments as Spectrum\_1D objects, the argument must follow ppm\_f1. As an example, to project the first experiment, one should type

```
s.projf2(1)
```

The user can replace this 'standard' numbering with the actual parameter that is varied during the evolution of the indirect dimension, by substitution of the ppm\_f1 attribute. As a result, the projection must be performed according to this new scale.

The phase adjustment is performed on a reference spectrum, then applied on the whole 2D matrix. By default, the chosen spectrum is the first one, but you can choose the one that fits the most your needs.

```
s.process()
s.pknl()  # Tries to remove the digital filter
s.adjph(expno = 10) # Calls interactive_phase_1D on the 10th experiment
```

The method plot shows the 2D contour map of the spectrum, just like the one of Spectrum\_2D. However, this is not always the most intelligent way to plot the data in order to gather information. This is the reason why this class features two unique additional methods that plot data: plot\_md and plot\_stacked. Both rely on the parameter which, that is a string of code (i.e. it should be interpreted by eval) that identifies which experiment to show by pointing at their index. which = 'all' results in pointing at all spectra.

```
s.plot() # 2D contour map
s.plot_md(which="3, 5, 11") # Plot the 3rd, the 5th and the 11th spectrum, superimposed
s.plot_stacked(which="np.arange(0,100,5)") # Makes a stacked plot with a spectrum every 5
```

The method integrate differs a little bit from the one coded in Spectrum\_1D.

```
s.integrate(which=2) # Interactive panel on the 3rd spectrum
```

Even if you select the integration limits on a single spectrum, the method integrate will compute

the integrals throughout the whole range of experiment. This means that each entry of integrals will be an array as long as the number of experiment.

### 2.6 Deconvolution of 1D datasets

The class fit. Voigt\_Fit in KLASSEZ offers a very convenient interface to deconvolve a spectrum by fitting. A shortcut to the class, which initializes the parameters automatically, is implemented in the attribute F of Spectrum\_1D.

To generate the input guess for the fit, you have to call the method iguess of the class. This can work in two different modes: the default one, which allows to build the guess peak-by-peak, and with auto=True, that features a peak-picker for the selection. The former is more precise, the second is much faster.

Whatever the employed method, the building of the initial guess is a two-stage process. First, you must zoom in with the matplotlib interactive viewer on the region of the spectrum you are interested in. Then, you can build the guess following the instructions in the GUI. When you press 'SAVE', your guess is stored, and the spectrum returns to the original view.

The 'manual' mode allows to optimize a polynomial baseline for each interval. A button labelled 'SET BASL' must be pressed when a satisfying region is highlighted in the GUI: this allows the scale on which the baseline is computed to be correctly aligned to the region itself. When this step is correctly performed, the box next to the button turns from red to green. Should the region be moved during the optimization of the initial guess, the box turns back to red, and the 'SET BASL' button must be pressed again to adjust the baseline scale accordingly.

The information on the peaks is saved in a .vf file, which can be imported with the function fit.read\_vf. There are two kind of .vf file: .ivf, that marks initial guesses, and .fvf, for the results of the fit. However, this is a human-only distinction, as the structure of the files is the same.

An example of .vf file is shown here:

```
! Initial guess computed by francesco on 11/11/2024 at 15:48:44
        Region;
                   Intensity
193.317:168.041; 8.08246575e+00
  #;
                           fwhm; Rel. I.;
                                             Phase;
                                                          Beta;
                                                                  Group
  1; 179.94060191; 172.500000; 1.0000000; -10.000; 0.000000;
        Region;
                 Intensity
   59.936:6.662; 5.02908980e+01
                           fwhm; Rel. I.;
                                              Phase:
  2;
       40.29851786;
                    150.000000; 0.214286;
                                               0.000;
                                                        0.00000;
                                                                       0
                      140.000000; 0.785714;
                                              10.000;
                                                        0.00000;
       24.98695246;
```

The header line, that starts with a !, is a comment, and acts as a separator between different attempts of the fit. In fact, .vf files are never overwritten: working again on the same file appends the information at the bottom. Hence, there is a parameter n in the fit.read\_vf function that allows to select which attempt to read.

Then, a series of blocks follow. Each block marks a region of selection: the keys 'Region' and 'Intensity' mark the limits of the fitting window, and the total intensity of the peaks. Under this line, there is a table that collects the peak parameters. As a final information there might be the baseline coefficients for the given region, which start with the key 'BASL\_C'. Should this line be missing, it means that the baseline was not optimized during the computation of the guess, and the coefficients will all be set to 0 when the file is read. The end of the block is marked with a line of '='

The method iguess automatically search for the existing input file. If it finds it, it is automatically loaded. Otherwise, the GUI for the computation of the initial guess opens up.

The fit can be performed by calling the method dofit, which returns a list of lmfit.MinimizerResult objects (one for each region) for a detailed inspection on how the fit performed. The behavior of the fit can be customized by setting the parameters of the method (see examples or the dedicated page of the manual). The fit goes region-by-region, and the results are saved in a .fvf file.

A .fvf file can be loaded using the method load\_fit.

Either the initial guess or the result of the fit can be conveniently visualized by using the method plot. Alternatively, the arrays of the model can be retrieved by calling calc\_fit\_lines. The method res\_histogram computes the histogram of the residuals, for a better understanding of the outcome of the fit procedure.

### 2.7 Example scripts

### 2.7.1 Reading and processing of 1D spectra

```
#! /usr/bin/env python3
from klassez import *
# Be aware that this is a BASIC processing
# Read the documentation of the functions to see the full powers
if 1:
    # This example is for the simulated data
   s = Spectrum_1D('acqus_1D', isexp=False)
   s.to_vf() # You can convert info on peaks to .ivf for fitting
else:
    # Use the following to read experimentals:
   spect = 'bruker', 'jeol', 'varian', 'magritek', 'oxford' # One of these
   s = Spectrum_1D(path_to_dataset, spect=spect)
# Setup the processing
   Apodization
       Follow the table in the user manual to see what reads what
s.procs['wf']['mode'] = 'em'
s.procs['wf']['lb'] = 5
   Zero-filling
s.procs['zf'] = 2**14
   Apply processing and do FT
```

```
s.process()
# Remove the digital filter
s.pknl()
# Phase correction
s.adjph()
# Plot the data
s.plot()
```

### 2.7.2 Fit 1D spectrum

The beginning of the script is the same of the reading example.

```
# s.F is a fit. Voiqt_Fit object
filename = 'test_1D_fit' # base filename for everything fit-related
# Compute the initial guess
auto = False
                 # True for peak-picker, False for manual
s.F.iguess(filename=filename, auto=auto)
if 0:
       # Do the fit
   lmfit_result = s.F.dofit( ### Parameters of the fitting ###
                           # movement for chemical shift /ppm
          u_lim=5,
          f_lim=50,
                           # movement for linewidth /Hz
          k_lim=(0, 3), # limits for intensity
          vary_phase=True, # optimize the phase of the peak
                            # optimize the lineshape (L/G ratio)
          vary_b=True,
          method='leastsq', # optimization method
          itermax=10000,
                            # max. number of iterations
          fit_tol=1e-10,
                            # arrest criterion threshold (see lmfit for details)
       basl_fit='fixed' # how to handle the baseline during the fit
          filename=filename, # filename for the .fvf file
else:
   # Load an existing .fuf file
   s.F.load_fit(filename=filename)
# Plot the results
s.F.plot(what='result', # what='iguess' for initial guess
        show_total=True, # Show the total trace or not
        show_res=True, \# Show the residuals
        res_offset=0.1, # Displacement of the residuals (plots residuals - res_offset)
                        # Labels for the peaks
        labels=None,
        filename=filename, # Filename for the figures
        ext='png', # format of the figure
        dpi=300,
                         # Resolution of the figure
# Compute histogram of the residuals
s.F.res_histogram(what='result',
            nbins=500,
                         # Number of bins of the histogram
            density=True, # Normalize them
            f_lims=None, # Limits for x axis
            xlabel='Residuals', # Guess what!
            x_symm=True, # Symmetrize the x-scale
            barcolor='tab:green', # Color of the bars
```

### 2.7.3 Read and process 2D spectrum

```
#! /usr/bin/env python3
from klassez import *
# Be aware that this is a BASIC processing
# Read the documentation of the functions to see the full powers
if 1:
   # This example is for the simulated data
   s = Spectrum_2D('acqus_2D', isexp=False)
    # For experimentals, at version 0.4a.7 klassez reads only 2D bruker
   s = Spectrum_2D(path_to_dataset)
# Setup the processing
   A podization
       Follow the table in the user manual to see what reads what
       REMEMBER: index 0 is F1, index 1 is F2, for procs
s.procs['wf'][1]['mode'] = 'em'
s.procs['wf'][1]['lb'] = 5
s.procs['wf'][0]['mode'] = 'qsin'
s.procs['wf'][0]['ssb'] = 2
   Zero-filling
s.procs['zf'] = 512, 2048
   Apply processing and do FT
s.process()
# Remove the digital filter
s.pknl()
# Phase correction
s.adjph()
# Plot the data
s.plot()
# Extract projections
ppm_f2 = 180
ppm_f1 = 10
s.projf1(ppm_f2) # Extract F1 trace @ ppm_f2 ppm
f1 = s.Trf1[f'{ppm_f2:.2f}'] # Call it back: it is a Spectrum_1D object!
f1.plot()
s.projf2(ppm_f1) # Extract F2 trace @ ppm_f1 ppm
f2 = s.Trf2[f'{ppm_f1:.2f}'] # Call it back: it is a Spectrum_1D object!
f2.plot()
```

# 3. List of modules and functions

# 3.1 MISC package

This package contains miscellaneous functions for the calculation of several properties, and generally for the handling of NMR spectra.

### $3.1.1 \quad misc.avg\_antidiag(X)$

Given a matrix X without any specific structure, finds the closest Hankel matrix in the Frobenius norm sense by averaging the antidiagonals.

#### Parameters:

• X: 2darray
Input matrix

#### Returns:

• Xp: 2darray
Hankel matrix obtained from X

# 3.1.2 misc.binomial\_triangle(n)

Calculates the n-th row of the binomial triangle. The first row is  $n=1,\,\mathrm{not}\ 0.$  Example:

### Parameters:

• n: *int* Row index

### **Returns:**

• row: *1darray*The n-th row of binomial triangle.

# 3.1.3 misc.calcres(fqscale)

Calculates the frequency resolution of an axis scale, i.e. how many Hz is a 'tick'.

### Parameters:

• fqscale : *1darray* Scale to be processed

### Returns:

ullet res: float The resolution of the scale

# $3.1.4 \quad misc.cmap2list(cmap, N=10, start=0, end=1)$

Extract the colors from a colormap and return it as a list.

#### Parameters:

- cmap: matplotlib.Colormap Object

  The colormap from which you want to extract the list of colors
- N: *int* Number of samples to extract
- start: float
  Start point of the sampling. 0 = beginning of the cmap; 1 = end of the cmap.
- end: float
  End point of the sampling. 0 = beginning of the cmap; 1 = end of the cmap.

#### Returns:

• colors: *list*List of the extracted colors.

### 3.1.5 misc.data2wav(data, filename='audiofile', cutoff=None, rate=44100)

Converts an array of data in a .wav file. The data are converted in float 32 format, then normalized to fit the (-1, 1) interval

#### Parameters:

• data: ndarray
Data to listen to

 $\bullet$  filename: str

Filename for the .wav file, without extension

• cutoff: *float or None* Clipping borders for the audio. If None, no clipping is performed

 $\bullet$  rate: int Sample rate in samples/sec

# $3.1.6 \quad misc.detect\_jumps(a)$

Detects where array a changes value.

### Parameters:

• a: 1darray Mask

### Returns:

• starts: *list*Left border of the windows /pt

• ends: *list*Right border of the windows /pt

### 3.1.7 misc.edit checkboxes(checkbox, xadj=0, yadj=0, dim=100, color=None)

Edit the size of the box to be checked, and adjust the lines accordingly.

#### **Parameters:**

- checkbox: matplotlib.widgets.CheckBox Object
  The checkbox to edit
- xadj: *float* modifier value for bottom left corner x-coordinate of the rectangle, in checkbox.ax coordinates
- yadj: *float* modifier value for bottom left corner y-coordinate of the rectangle, in checkbox.ax coordinates
- dim: *float*Area of the square, in pixels. Default value is 25
- color: str or list or None
  If it is not None, change color to the lines

# $3.1.8 \quad misc.extend\_taq(old\_taq, newsize=None)$

Extend the acquisition timescale to a longer size, using the same dwell time

### Parameters:

- old\_taq: 1darray Old timescale
- ullet new size: int New size of acquisition timescale, in points

### **Returns:**

• new\_taq: *1darray* Extended timescale

# 3.1.9 misc.find\_nearest(array, value)

Finds the value in array which is the nearest to value .

### Parameters:

• array : *1darray* Self-explanatory

ullet value : float

Value to be found

### Returns:

• val : *float*The closest value in array tovalue

# $3.1.10 \quad misc.freq2ppm(x, B0=701.125, o1p=0)$

Converts xfrom Hz to ppm.

### Parameters:

• x : float Value to be converted

• B0 : float Field frequency, in MHz. Default: 700 MHz

• o1p : float Carrier frequency, in ppm. Default: 0.

### Returns:

• y : *float*The converted value

### 3.1.11 misc.get trace(data, ppm f2, ppm f1, a, b=None, column=True)

Takes as input a 2D dataset and the ppm scales of direct and indirect dimensions respectively. Calculates the projection on the given axis summing from a (ppm) to b (ppm). Default: indirect dimension projection (i.e. column=True), change it to 'False' for the direct dimension projection.

#### Parameters:

- data : 2darray

  Spectrum of which to extract the projections
- ppm\_f2: 1darray ppm scale of the direct dimension
- ppm\_f1 : *1darray* ppm scale of the indirect dimension
- a: float

  The ppm value from which to start extracting the projection.
- b: float, optional
  If provided, the ppm value at which to stop extracting the projection. Otherwise, returns only the 'a' trace.
- column: bool

  If True, extracts the F1 projection. If False, extracts the F2 projection.

#### Returns:

• y : *1darray*Computed projection

# 3.1.12 misc.get\_ylim(data\_inp)

Calculates the y-limits of ax as follows:

- Bottom: min(data) 5% max(height)
- Top: max(data) + 5% max(height)

where height = max(data) - min(data)

### Parameters:

• data\_inp: ndarray or list Input data. If it is a list, data\_inp is converted to array.

### Returns:

• lims: tuple Bottom, Top

### 3.1.13 misc.hankel(data, n=None)

Computes a Hankel matrix from data. If data is a 1darray of length N, computes the correspondent Hankel matrix of dimensions (N-n+1, n). If data id a 2darray, computes the closest Hankel matrix in the Frobenius norm sense by averaging the values on the antidiagonals.

### Parameters:

- data: *1darray* Vector to be Hankel-ized, of length N
- $\bullet$  n: intNumber of columns that the Hankel matrix will have

#### Returns:

• H: 2darray
Hankel matrix of dimensions (N-n+1, n)

# 3.1.14 misc.hz2pt(fqscale, hz)

Converts hz from frequency units to points, on the basis of its scale.

### Parameters:

• fqscale : *1darray* Scale to be processed

• hz : *float* Value to be converted

### Returns:

• pt : *float*The frequency value converted in points

# 3.1.15 misc.in2px(\*in\_args)

Converts a sequence of numbers from inches to pixels by multiplying times 96.

### Parameters:

• \*in\_args: sequence of floats Values in inches to convert

### Returns:

• px\_args: tuple of ints Values in pixels

# 3.1.16 misc.lenslice(a)

Calculates the length of a slice, i.e. the length an array would have when sliced with this slice.

### Parameters:

ullet a: slice Slice of which to calculate the length

### Returns:

• length: *int* Length of the slice

### 3.1.17 misc.load ser(path, TD1=1, BYTORDA=0, DTYPA=0, cplx=True)

Reads a binary file and transforms it in an array. The parameters BYTORDA and DTYPA can be found in the acqus file.

- BYTORDA =  $1 \rightarrow \text{big endian} \rightarrow '>'$
- BYTORDA =  $0 \rightarrow \text{little endian} \rightarrow '<'$
- DTYPA =  $0 \rightarrow \text{int} 32 \rightarrow 'i4'$
- DTYPA =  $2 \rightarrow \text{float}64 \rightarrow \text{'f8'}$

#### **Parameters:**

- path : strPath to the file to read
- TD1: *int*Number of experiments in the indirect dimension
- BYTORDA: int Endianness of data
- DTYPA: *int*Data type format
- cplx: bool

  If True, the input data are interpreted as complex, which means that in the direct dimension there will be real and imaginary parts alternated.

#### Returns:

• data: 2darray Array of data.

# 3.1.18 misc.makeacqus 1D(dic)

Given a NMRGLUE dictionary from a 1D spectrum (generated by ng.bruker.read), this function builds the acqus file with only the 'important' parameters.

### Parameters:

 $\bullet$  dic: dict NMRglue dictionary returned by ng.bruker.read

### Returns:

# 3.1.19 misc.makeacqus\_1D\_jeol(dic)

Given a dictionary from a 1D spectrum (generated by jeol\_parser.parse), this function builds the acqus file with only the 'important' parameters.

### Parameters:

### Returns:

### 3.1.20 misc.makeacqus 1D oxford(dic)

Given a NMRGLUE dictionary from a 1D spectrum (generated by ng.jcampdx.read), this function builds the acqus file with only the 'important' parameters.

### Parameters:

### Returns:

### 3.1.21 misc.makeacqus 1D spinsolve(dic)

Given a NMRGLUE dictionary from a 1D spectrum (generated by ng.spinsolve.read), this function builds the acqus file with only the 'important' parameters. Be sure to get the info from all the configuration files!

### Parameters:

• dic: dict

NMRglue dictionary returned by ng.spinsolve.read

### Returns:

### 3.1.22 misc.makeacqus 1D varian(dic)

Given a NMRGLUE dictionary from a 1D spectrum (generated by ng.varian.read), this function builds the acqus file with only the 'important' parameters.

### Parameters:

### Returns:

# 3.1.23 misc.makeacqus 2D(dic)

Given a NMRGLUE dictionary from a 2D spectrum (generated by ng.bruker.read ), this function builds the acqus file with only the 'important' parameters.

### Parameters:

 $\bullet$  dic: dict NMRglue dictionary returned by ng.bruker.read

### Returns:

### 3.1.24 misc.mathformat(ax, axis='y', limits=(-2, 2))

Apply exponential formatting to the given axis of the given figure panel. The offset text size is uniformed to the tick labels' size.

- ax: matplotlib.Subplot Object Panel of the figure to edit
- axis: *str* 'x', 'y' or 'both'.
- limits: *tuple* tuple of ints that indicate the order of magnitude range outside which the exponential format is applied.

# 3.1.25 misc.merge\_dict(\*dics)

Merge a sequence of dictionaries in a single dictionary.

### Parameters:

• dics: sequence of dict Dictionaries to merge

### Returns:

 $\bullet \ \, \mathrm{merged\_dict} \colon \, dict \\ \mathrm{Merged\ dictionary}$ 

# 3.1.26 misc.molfrac(n)

Computes the 'molar fraction' 'x' of the array 'n'. Also computes the total amount.

### Parameters:

• n: list or 1darray list of values

### Returns:

- x: *list or 1darray* molar fraction array
- N: *float* sum of all the elements in 'n'

### 3.1.27 misc.noise std(y)

Calculates the standard deviation of the noise using the Bruker formula. Taken y as an array of r points, and y[k] its k-th entry:

$$\sigma_N = \frac{1}{\sqrt{r-1}} \sqrt{\sum_{k=0}^{r-1} (\mathbf{y}[k]^2) - \frac{1}{r} \left[ \left( \sum_{k=0}^{r-1} \mathbf{y}[k] \right)^2 + \frac{3}{r^2-1} \left( \sum_{k=0}^{\lfloor r/2 \rfloor - 1} (k+1) (\mathbf{y}[\lfloor r/2 \rfloor + k] - \mathbf{y}[\lfloor r/2 \rfloor - k - 1]) \right)^2 \right]}$$

### Parameters:

 $\bullet$  y : 1darray The spectral region you would like to use to calculate the standard deviation of the noise.

#### Returns:

• noisestd : *float*The standard deviation of the noise.

# 3.1.28 misc.nuc\_format(nuc)

Converts the 'nuc' key you may find in acqus in the formatted label, e.g. '13C'  $\rightarrow$  '\$^{13}\$C'

### Parameters:

 $\bullet$  nuc: str Unformatted string

### Returns:

 $\bullet$  fnuc: str Formatted string.

# 3.1.29 misc.polyn(x, c)

Computes p(x), polynomion of degree n-1, where n is the number of provided coefficients.

#### Parameters:

- $\bullet$  x : 1darray Scale upon which to build the polynomion
- c : list or 1darray Sequence of the polynomion coefficient, starting from the 0-th order coefficient

### Returns:

• px : *1darray* Polynomion of degree n-1.

# $3.1.30 \quad misc.ppm2freq(x, B0=701.125, o1p=0)$

Converts xfrom ppm to Hz.

### Parameters:

• x : float Value to be converted

• B0 : float Field frequency, in MHz. Default: 700 MHz

• o1p : float Carrier frequency, in ppm. Default: 0.

### Returns:

• y : *float*The converted value

# 3.1.31 misc.ppmfind(ppm\_scale, value)

Finds the exact value in ppm\_scale.

### Parameters:

• ppm\_scale : 1darray Self-explanatory

ullet value : float

The value to be found

### Returns:

• I : int
The index correspondant to 'V' in 'ppm\_scale'

• V : float
The closest value to 'value' in 'ppm\_scale'

# 3.1.32 misc.pretty\_scale(ax, limits, axis='x', n\_major\_ticks=10, \*, minor\_each=5, fmt=None)

This function computes a pretty scale for your plot. Calculates and sets a scale made of 'n\_major\_ticks' numbered ticks, spaced by 'minor\_each' unnumbered ticks. After that, the plot borders are trimmed according to the given limits.

- ax: matplotlib.AxesSubplot object
  Panel of the figure of which to calculate the scale
- limits: *tuple* limits to apply of the given axis. (left, right)
- axis: str'x' for x-axis, 'y' for y-axis, 'z' for z-axis
- n\_major\_ticks: *int*Number of numbered ticks in the final scale. An oculated choice gives very pleasant results.
- minor\_each: *int*Number of divisions for each interval between two major ticks
- fmt: strString-formatting for the numbers on the axis. Should be given as e.g. '.3f'

# $3.1.33 \quad misc.print\_dict(mydict)$

Prints a dictionary one entry per row, in the format key: value. Nested dictionaries are printed with an indentation

### Parameters:

• mydict: *dict*The dictionary you want to print

### Returns:

ullet outstring: strThe printed text formatted as single string

# $3.1.34 \quad misc.print\_list(mylist)$

Prints a list, one entry per row.

### Parameters:

 $\bullet$  mylist: listThe list you want to print

### Returns:

ullet outstring: strThe printed text formatted as single string

# 3.1.35 misc.procpar(txt)

Takes as input the path of a file containing a 'key' in the first column and a 'value' in the second column. Returns the correspondant dictionary

### Parameters:

 $\bullet$  txt : strPath to a file that contains 'key' in first column and 'value' in the second

### Returns:

• procpars : dict
Dictionary of shape 'key':'value'

# $3.1.36 \quad misc.px2in(*px\_args)$

Converts a sequence of numbers from inches to pixels by multiplying times 96.

### Parameters:

• \*px\_args: sequence of ints Values in pixels to convert

### Returns:

• in\_args: tuple of floats Values in inches

### 3.1.37 misc.readlistfile(datafile)

Takes as input the path of a file containing one entry for each row. Returns a list of the aforementioned entries.

### Parameters:

ullet datafile: strPath to a file that contains one entry for each row

### Returns:

• files: *list*List of the entries contained in the file

### 3.1.38 misc.select for integration(ppm f1, ppm f2, data, Neg=True)

Select the peaks of a 2D spectrum to integrate. First, select the area where your peak is located by dragging the red square. Then, select the center of the peak by right\_clicking. Finally, click 'ADD' to store the peak. Repeat the procedure for as many peaks as you want.

### Parameters:

• ppm\_f1 : *1darray* ppm scale of the indirect dimension

• ppm\_f2 : *1darray* ppm scale of the direct dimension

• data : 2darray Spectrum

• Neg: bool Choose if to show the negative contours ( True) or not ( False )

#### Returns:

• peaks: list of dict
For each peak there are two keys, 'f1' and 'f2', whose meaning is obvious. For each of these keys, you have 'u': center of the peak /ppm, and 'lim': the limits of the square you drew before.

### 3.1.39 misc.select traces(ppm f1, ppm f2, data, Neg=True, grid=False)

Select traces from a 2D spectrum, save the coordinates in a list. Left click to select a point, right click to remove it.

### Parameters:

- ppm\_f1 : *1darray* ppm scale of the indirect dimension
- ppm\_f2 : *1darray* ppm scale of the direct dimension
- data : *2darray* Spectrum
- Neg: bool Choose if to show the negative contours ( True) or not ( False )
- grid : bool Choose if to display the grid ( True) or not ( False )

### Returns:

• coord: *list*List containing the '[x,y]' coordinates of the selected points.

### 3.1.40 misc.set fontsizes(ax, fontsize=10)

Automatically adjusts the fontsizes of all the figure elements. In particular:

- $\bullet$  title = fontsize
- axis labels = fontsize 2
- ticks labels = fontsize 3
- $\bullet$  legend entries = fontsize 4

- ax: matplotlib.Subplot Object Subplot of interest
- fontsize: *float* Starting fontsize

# 3.1.41 misc.set\_ylim(ax, data\_inp)

Set the limits on the y-axis on the ax subplot. The values are computed using misc.get\_ylim.

- ax: matplotlib.Subplot Object
  Panel of the figure where to apply this scale
- data\_inp: ndarray or list Input data. If it is a list, data\_inp is converted to array.

### 3.1.42 misc.show cmap(cmap, N=10, start=0, end=1, filename=None)

Plot the colors extracted from a colormap.

- cmap: matplotlib.Colormap Object

  The colormap from which you want to extract the list of colors
- N: *int* Number of samples to extract
- start: float
  Start point of the sampling. 0 = beginning of the cmap; 1 = end of the cmap.
- end: float
  End point of the sampling. 0 = beginning of the cmap; 1 = end of the cmap.
- filename:  $str\ or\ None$  Filename of the figure to be saved. The '.png' extension is added automatically. If None, the figure is shown instead

### 3.1.43 misc.snr(data, signal=None, n reg=None)

Computes the signal to noise ratio of a 1D spectrum.

#### Parameters:

- $\bullet$  data : 1darray The spectrum of which you want to compute the SNR
- signal : float, optional

  If provided, uses this value as maximum signal. Otherwise, it is selected as the maximum value in 'data'
- n\_reg: list or tuple, optional If provided, contains the points that delimit the noise region. Otherwise, the whole spectrum is used.

#### Returns:

• snr : *float* The SNR of the spectrum

### 3.1.44 misc.snr 2D(data, n reg=None)

Computes the signal to noise ratio of a 2D spectrum.

#### Parameters:

 $\bullet$  data : 1darray The spectrum of which you want to compute the SNR

• n\_reg: list or tuple
If provided, the points of F1 scale and F2 scale, respectively, of which to extract the projections.
Otherwise, opens the tool for interactive selection.

#### Returns:

• snr\_f1 : float
The SNR of the indirect dimension

• snr\_f2 : float
The SNR of the direct dimension

# 3.1.45 misc.split\_acqus\_2D(acqus)

Split the acqus dictionary of a 2D spectrum into two separate 1D-like acqus dictionaries.

### Parameters:

ullet acqus: dict acqus dictionary of a 2D spectrum

### Returns:

- acqu1s: *dict* acqus dictionary of the indirect dimension
- acqu2s: *dict* acqus dictionary of the direct dimension

# $3.1.46 \quad misc.split\_procs\_2D(procs)$

Split the procs dictionary of a 2D spectrum into two separate 1D-like procs dictionaries.

### Parameters:

ullet procs: dict procs dictionary of a 2D spectrum

### Returns:

• proc1s: *dict* procs dictionary of the indirect dimension

• proc2s: *dict* procs dictionary of the direct dimension

# 3.1.47 misc.sum\_overlay(y1, y2, x0, x=None)

Compute the sum of two arrays  $y_1$  and  $y_2$  of different dimensions. The sum starts at  $x_0$  (on the scale x) as left anchor point.

#### Parameters:

- y1: *1darray* Original array
- y2: 1darray Array to sum
- x0: float Value on the x scale as left anchor point
- x: 1darray
  Reference scale. If None, the index points is used.

#### Returns:

• ym: *1darray* Summed arrays

# 3.1.48 misc.trim\_data(ppm\_scale, y, lims)

Trims the frequency scale and correspondant 1D dataset y from sx (ppm) to dx (ppm).

#### Parameters:

- ppm\_scale : 1darray ppm scale of the spectrum
- y : 1darray spectrum
- lims: *tuple* ppm values where to start and stop trimming

#### Returns:

 $\bullet$  xtrim : 1darray Trimmed ppm scale

 $\bullet$  ytrim : 1darray Trimmed spectrum

# $3.1.49 \quad misc.trim\_data\_2D(x\_scale, y\_scale, data, xlim=None, ylim=None)$

Trims data and the scales according to xlim and ylim. Returns the trimmed data and the correspondant trimmed scales.

#### Parameters:

- x\_scale: 1darray Scale for the rows of data
- y\_scale: *1darray* Scale for the columns of data
- data: *2darray*Data to be trimmed
- xlim: tuple Limits for x\_scale (L, R)
- ylim: tuple Limits for y\_scale (L, R)

#### Returns:

- trimmed\_x: 1darray
  Trimmed x scale
- trimmed\_y: 1darray
  Trimmed y\_scale
- trimmed\_data: 2darray Trimmed data

# 3.1.50 misc.unhankel(H)

Concatenates the first row and the last column of the matrix H, which should have Hankel-like structure, so to build the array of independent parameters.

#### Parameters:

• H: 2darray Hankel-like matrix

#### Returns:

• h: *1darray*First row and last column, concatenated

# 3.1.51 misc.write\_acqus\_1D(acqus, path='sim\_in\_1D')

Writes the input file for a simulated spectrum, basing on a dictionary of parameters.

#### Parameters:

 $\bullet$  acqus : dict The dictionary containing the parameters for the simulation

• path: str, optional
Directory where the file will be saved.

# 3.1.52 misc.write\_acqus\_2D(acqus, path='sim\_in\_2D')

Writes the input file for a simulated spectrum, basing on a dictionary of parameters.

#### Parameters:

 $\bullet$  acqus : dict The dictionary containing the parameters for the simulation

• path: str, optional
Directory where the file will be saved.

# 3.1.53 misc.write\_help(request, file=None)

Gets the documentation of request, and tries to save it in a text file.

#### Parameters:

- request: function or class or package Whatever you need documentation of
- file: str or None or False

  Name of the output documentation file. If it is None, a default name is given. If it is False, the output is printed on screen.

# 3.1.54 misc.write\_ser(fid, path='./', BYTORDA=0, DTYPA=0, overwrite=True)

Writes the FID file in directory 'path', in a TopSpin-readable way (i.e. little endian, int32). The binary file is named 'fid' if 1D, 'ser' if multiD.

- BYTORDA =  $1 \rightarrow \text{big endian} \rightarrow '>'$
- BYTORDA =  $0 \rightarrow \text{little endian} \rightarrow '<'$
- DTYPA =  $0 \rightarrow \text{int}32 \rightarrow 'i4'$
- DTYPA =  $2 \rightarrow \text{float64} \rightarrow \text{'f8'}$

#### Parameters:

- fid : *ndarray* FID array to be written
- path : strDirectory where to save the file

# 3.1.55 misc.zero crossing(array, after=False)

Find the indices where the elements in the array change sign. The identified positions are the ones before the sign changes. This behavior can be modified by setting 'after=True'.

#### Parameters:

• array: *1darray*Data to analyze

• after: bool

If True, returns the indices of the element after the sign change; if False, the indices before.

#### Returns:

• zerocross: *1darray*Position of the zero-crossing, according to 'before'

# 3.2 PROCESSING package

This package contains functions for the processing of NMR spectra, either in time domain or in frequency domain, and the transition between the two domains.

# 3.2.1 processing.abc(ppm, data, n=5, lims=None, alpha=2.75, qfil=False, qfilp={'u': 4.7, 's': 10})

Automatic computation of a baseline for a spectrum using a thresholding-based method for the detection of the baseline-only region, followed by a weighted linear least squares optimization with a polynomion of degree n-1. The weights are computed on the absolute value of the first derivative of the spectrum. Set qfil to True if there is a very intense solvent peak that would hamper the computation of the threshold.

#### Parameters:

- ppm: *1darray* PPM scale of the spectrum
- data: *1darray*The spectrum to baseline-correct
- n: *int*Number of coefficients of the polynomial baseline
- lims: tuple or None
  Limits for the region on which to compute the baseline, in ppm
- alpha: float
  The threshold will be set as thr = alpha \* np.std(np.gradient(data))
- qfil: bool
  Choose whether to apply a filter on the solvent region (True) or not (False)
- qfilp: dict
  - 'u' = center of the filter in ppm
  - 's' = width of the filter in Hz

#### Returns:

• baseline: *1darray*Computed baseline

# 3.2.2 processing.abs(ppm, data, n=5, lims=None, alpha=2.75, qfil=False, qfilp= $\{'u': 4.7, 's': 10\}$ )

Computes the baseline correction on data using processing.abc, and gives back the subtracted spectrum. The imaginary part of the spectrum is reconstructed using processing.hilbert.

#### Parameters:

• ppm: 1darray

PPM scale of the spectrum

• data: 1darray

The spectrum to baseline-correct

• n: *int* 

Number of coefficients of the polynomial baseline

• lims: tuple or None

Limits for the region on which to compute the baseline, in ppm

• alpha: float

The threshold will be set as thr = alpha \* np.std(np.gradient(data))

• qfil: bool

Choose whether to apply a filter on the solvent region (True) or not (False)

 $\bullet$  qfilp: dict

- 'u' = center of the filter in ppm

- 's' = width of the filter in Hz

#### Returns:

• S: 1darray
Baseline-subtracted spectrum

### 3.2.3 processing.acme(data, m=1, a=5e-05)

Automated phase Correction based on Minimization of Entropy. This algorithm allows for automatic phase correction by minimizing the entropy of the m-th derivative of the spectrum, as explained in detail by L. Chen et.al. in Journal of Magnetic Resonance 158 (2002) 164-168.

Defined the entropy of  $\mathbb{h}$  as:

$$S = -\sum_{j} \mathbb{h}[j] \ln(\mathbb{h}[j])$$

and

$$h = \frac{\left| \mathbb{r}[j]^{(m)} \right|}{\sum_{j} \left| \mathbb{r}[j]^{(m)} \right|}$$

where

$$\mathbb{r} = \operatorname{Re}\{\operatorname{spectrum} e^{-\mathrm{i}\phi}\}$$

and  $\mathbf{r}^{(m)}$  is the *m*-th derivative of  $\mathbf{r}$ , the objective function to minimize is:

$$S + P(\mathbf{r})$$

where  $P(\mathbf{r})$  is a penalty function for negative values of the spectrum.

The phase correction is applied using processing.ps. The values p0 and p1 are fitted using Nelder-Mead algorithm.

#### **Parameters:**

- data: *1darray*Spectrum to be phased, complex
- m: *int*Order of the derivative to be computed
- a: *float*Weighting factor for the penalty function

#### Returns:

- p0f: float
  Fitted zero-order phase correction, in degrees
- p1f: *float*Fitted first-order phase correction, in degrees

## 3.2.4 processing.align(ppm scale, data, lims, u off=0.5, ref idx=0)

Performs the calibration of a pseudo-2D experiment by circular-shifting the spectra of an appropriate amount. The target function aims to minimize the superimposition between a reference spectrum and the others using a brute-force method.

#### Parameters:

- ppm\_scale: *1darray* ppm scale of the spectrum to calibrate
- data: *2darray*Complex-valued spectrum
- lims: tuple (ppm sx, ppm dx) of the calibration region
- u\_off: float
  Maximum offset for the circular shift, in ppm
- ref\_idx: *int*Index of the spectrum to be used as reference

#### Returns:

- data\_roll: 2darray Calibrated data
- u\_cal: *list*Number of point of which the spectra have been circular-shifted
- u\_cal\_ppm: *list*Correction for the ppm scale of each experiment

# 3.2.5 processing.apk(ppm, data, SFO1, alpha=3, winsize=50, ap1=True, seethrough=False)

Performs automatic phase correction.

The algorithm starts with the computation of a mask to separate signal from baseline-only regions. This is done via an iterative thresholding, i.e. a point is 'signal' if the first derivative of the spectrum in that point is higher than its standard deviation by  $\alpha$  times:

$$ifd'[k] > \alpha\sigma(d') \implies k \in signal region$$

The selection is further refined by repeating the same procedure on the original data. Then, the regions separated by less than winsize are joined together, and the presence of actual peaks in the region is checked with a pick-picker.

At this point, each region is phased independently with only phase 0. The phase angle is tested in a brute-force manner. The cost function minimizes the area below the straight line that connects the borders of the window. The first-order phase correction is calculated with a weighted linear regression, where the weights are the integrals of the magnitude of each region.

The choice of alpha and winsize can be important for the outcome. Higher alpha values make the detection of peaks more stringent -> for spectra with high SNR and sharp peaks a suitable value is 4-6. Decreasing winsize makes the algorithm to estimate more regions.

#### **Parameters:**

• ppm: *1darray* ppm scale of the spectrum

• data: *1darray* Spectrum

• alpha: float

• SFO1: float Nucleus' Larmor frequency /MHz

Nucleus' Larmor frequency / MHz

• winsize: *float*Minimum size of the window that can contain peaks /Hz

• ap1: bool

True to adjust both zero and first order, False for only phase zero

Factor that multiplies the std of the spectrum to set the threshold

• seethrough: bool

If True, draws a series of diagnostic figures to see what the algorithm is doing

#### Returns:

• datap: *1darray* Phased data

• values: *tuple*Found values (p0, p1)

# 3.2.6 processing.baseline\_correction(ppm, data, basl\_file='spectrum.basl', winlim=None)

Interactively corrects the baseline of a given spectrum and saves the parameters in a file. The program starts with an interface to partition the spectrum in windows to correct separately. Then, for each window, an interactive panel opens to allow the user to compute the baseline.

#### Parameters:

• ppm: *1darray* PPM scale of the spectrum

• data: *1darray*The spectrum of which to adjust the baseline

• basl\_file: str Name for the baseline parameters file

• winlim: list or str or None
List of the breakpoints for the window. If it is str, indicates the location of a file to be read with np.loadtxt. If it is None, the partitioning is done interactively.

# 3.2.7 processing.blp(data, pred=1, order=8)

Applies backward linear prediction by calling processing.lp with mode='b'.

#### Parameters:

• data: *1darray* FID to be linear-predicted

• pred: *int* Number of points to predict

ullet order: int Number of coefficients to use for the prediction

#### Returns:

• lpdata: *1darray* FID with linear prediction applied.

### 3.2.8 processing.blp ng(data, pred=1, order=8, N=2048)

Performs backwards linear prediction on data. This function calls nmrglue.process.proc\_lp.lp with most of the parameters set automatically. The algorithm predicts 'pred' points of the FID using 'order' coefficient for the linear interpolation. Only the first N points of the FID are used in the LP equation, because the computational cost scales with n\*\*2, making the use of more than 8k points not effective: using more points brings negligible contiribution to the final result. For Oxford spectra, set 'pred' to half the value written in 'TDoff'.

#### Parameters:

• data: *ndarray*Data on which to perform the linear prediction. For 2d data, it is performed row-by-row

• pred: *int*Number of points to be predicted

• order: *int*Number of coefficients to be used for the prediction

• N: *int*Number of points of the FID to be used in the calculation

#### Returns:

• datap: *ndarray*Data with the predicted points appended at the beginning

## 3.2.9 processing.cadzow(data, n, nc, print head=True)

This functions performs Cadzow denoising on  $\mathtt{data}$ , which is a 1D array of N points. The algorithm works as follows:

- 1. Transform data in a Hankel matrix  $\mathbb{H}$  of dimensions (N-n,n)
- 2. Make SVD on  $\mathbb{H} = \mathbb{USV}^H$
- 3. Keep only the first nc singular values, and put all the rest to 0 ( $\mathbb{S} \to \mathbb{S}'$ )
- 4. Rebuild  $\mathbb{H}' = \mathbb{US}'\mathbb{V}^H$
- 5. Average the antidiagonals to rebuild the Hankel-type structure, then make 1D array

#### **Parameters**

- data: *1darray* Input data
- n: *int*Number of columns of the Hankel matrix.
- nc: *int*Number of singular values to keep.
- print\_head: *bool*Set it to True to display the fancy heading.

#### Returns

• datap: *1darray* Denoised data

# $\begin{array}{ll} 3.2.10 & processing.cadzow\_2D(data,n,nc,i=True,itermax=100,f=0.005,\\ & print\_time=True) \end{array}$

Performs the Cadzow denoising method on a 2D spectrum, one transient at the time. This function calls cadzow if i=False, or iterCadzow if i=True.

#### **Parameters**

- data: 2darray Input data
- n: *int*Number of columns of the Hankel matrix.
- nc: *int* Number of singular values to keep.
- i: bool

  Calls processing.cadzow if i=False, or processing.iterCadzow if i=True.
- itermax: *int*Maximum number of iterations allowed.
- f: float
  Factor for the arrest criterion.
- print\_time: *bool*Set it to True to display the time spent.

#### Returns

• datap: 2darray
Denoised data

# 3.2.11 processing.calc\_nc(data, s\_n)

Calculates the optimal number of components, given the standard deviation of the noise. The threshold value is calculated as stated in Theorem 1 of reference: https://arxiv.org/abs/1710.09787v2

#### Parameters:

• data: *2darray* Input data

• s\_n: *float*Noise standard deviation

#### Returns:

• n\_c: *int* Number of components

### 3.2.12 processing.calibration(ppmscale, S)

Scroll the ppm scale of spectrum to make calibration. The interface offers two guidelines: the red one, labelled 'reference signal' remains fixed, whereas the green one ('calibration value') moves with the ppm scale. The ideal calibration procedure consists in placing the red line on the signal you want to use as reference, and the green line on the ppm value that the reference signal must assume in the calibrated spectrum. Then, scroll with the mouse until the two lines are superimposed.

#### Parameters:

- ppmscale: *1darray*The ppm scale to be calibrated
- S: *1darray*The spectrum to calibrate

#### Returns:

• offset: float
Difference between original scale and new scale. This must be summed up to the original ppm
scale to calibrate the spectrum.

### 3.2.13 processing.convdta(data, grpdly=0, scaling=1)

Removes the digital filtering to obtain a spectrum similar to the command CONVDTA performed by TopSpin. However, they will differ a little bit because of the digitization. These differences are not invisible to human's eye.

#### Parameters:

• data: *ndarray* FID with digital filter

• grpdly: int Number of points that the digital filter consists of. Key \$GRPDLY in acqus file

• scaling: float
Scaling factor of the resulting FID. Needed to match TopSpin's intensities.

#### Returns:

• data\_in: ndarray FID without the digital filter. It will have grpdly points less than data.

# 3.2.14 processing.convolve(in1, in2)

Perform the convolution of the two array by multiplying their inverse Fourier transform. The two arrays must have the same dimension.

#### Parameters:

• in1: ndarray
First array

• in2: ndarray
Second array

#### Returns:

• cnv: ndarray
Convolved array

# 3.2.15 processing.eae(data)

Shuffles data if the spectrum is acquired with FnMODE = Echo-Antiecho. NOTE: introduces -90° phase shift in F1, to be corrected after the processing

```
pdata = np.zeros_like(data)
pdata[::2] = (data[::2].real - data[1::2].real) + 1j*(data[::2].imag - data[1::2].imag)
pdata[1::2] = -(data[::2].imag + data[1::2].imag) + 1j*(data[::2].real + data[1::2].real)
```

# 3.2.16 processing.em(data, lb, sw)

Exponential apodization

#### Parameters:

- data: ndarray Input data
- lb: *float*Lorentzian broadening. It should be positive.

## 3.2.17 processing.fp(data, wf=None, zf=None, fcor=0.5, tdeff=0)

Performs the full processing of a 1D NMR FID (data).

#### Parameters:

- data: *1darray* Input data
- wf: dict {'mode': function to be used, 'parameters': different from each function}
- zf: *int* final size of spectrum
- fcor: *float* weighting factor for the FID first point
- tdeff: *int* number of points of the FID to be used for the processing.

#### Returns:

• datap: *1darray* Processed data

## 3.2.18 processing.ft(data0, alt=False, fcor=0.5)

Fourier transform in NMR sense. This means it returns the reversed spectrum.

#### Parameters:

- data0: ndarray Array to Fourier-transform
- alt: bool negates the sign of the odd points, then take their complex conjugate. Required for States-TPPI processing.
- fcor: float weighting factor for FID 1st point. Default value (0.5) prevents baseline offset

#### Returns:

• dataft: *ndarray* Transformed data

# 3.2.19 processing.gm(data, lb, gb, gc, sw)

Gaussian apodization. The parameter 'lb' controls the sharpening factor of a rising exponential, and behaves exactly as in processing.em. In contrast, 'gb' controls the gaussian decay factor. Apply this function VERY CAREFULLY. Choose the right values through the interactive processing.

#### Parameters:

- data: ndarray
  Input data
- lb: *float*Lorentzian sharpening /Hz. It should be negative.
- gb: float
  Gaussian broadening. It should be positive.
- gc: float Gaussian center, relatively to the FID length:  $0 \le gc \le 1$
- sw: *float* Spectral width /Hz

#### Returns:

• pdata: ndarray Processed data

# 3.2.20 processing.gmb(data, lb, gb, sw)

Bruker-style Gaussian apodization. Apply this function VERY CAREFULLY. Choose the right values through the interactive processing.

#### Parameters:

- data: ndarray Input data
- lb: *float* Lorentzian sharpening /Hz. It should be negative.
- gb: *float*Gaussian broadening. It should be positive.
- sw: float Spectral width /Hz

#### Returns:

• pdata: ndarray Processed data

## 3.2.21 processing.hilbert(f)

Computes the Hilbert transform of real vector f in order to retrieve its imaginary part. Make sure that the original spectrum was zero-filled to at least twice the original size of the FID. The algorithm computes the convolution by means of FT, as follows:

- make IFT of f = a
- compute h = [1j for x in range(N)if x<N/2 else -1j]
- Compute b = ha
- Build d = a + ib
- make FT of d = F
- replace Re(F) with f

#### Parameters:

• f: ndarray
Array of which you want to compute the imaginary part

#### Returns:

• f\_cplx: ndarray Complex version of f

# 3.2.22 processing.hilbert2(data)

Retrieve the imaginary parts of a hypercomplex dataset, when you only have the rr part.

- $rr = Re(Ht\{rr\})$
- $ir = Im(Ht\{rr\})$
- $ri = -Im(Ht\{rr.T\}.T)$
- $ii = Im(Ht\{ri\})$

#### Parameters:

• data: *2darray* rr part

#### Returns:

- rr: 2darray
  Real part in f2, real part in f1
- ir: 2darray
  Imaginary part in f2, real part in f1
- ri: 2darray
  Real part in f2, imaginary part in f1
- ii: 2darray Imaginary part in f2, imaginary part in f1

### 3.2.23 processing.ift(data0, alt=False, fcor=0.5)

Inverse Fourier transform in NMR sense. This means that the input dataset is reversed before to do iFT.

#### Parameters:

- data0: ndarray Array to Fourier-transform
- alt: bool negates the sign of the odd points, then take their complex conjugate. Required for States-TPPI processing.
- fcor: float weighting factor for FID 1st point. Default value (0.5) prevents baseline offset

#### Returns:

• dataft: ndarray
Transformed data

## 3.2.24 processing.integral(fx, x=None, lims=None)

Calculates the primitive of fx. If fx is a multidimensional array, the integrals are computed along the last dimension.

#### Parameters:

- fx: ndarray
  Function (array) to integrate
- x: 1darray or None Independent variable. Determines the integration step. If None, it is the point scale
- lims: tuple or None Integration range. If None, the whole function is integrated.

#### Returns:

• Fx: *ndarray* Integrated function.

#### 

Calculate the integral of a 2D peak. The idea is to extract the traces correspondent to the peak center and fit them with a gaussian function in each dimension. Then, once got the intensity of each of the two gaussians, multiply them together in order to obtain the 2D integral. This procedure should be equivalent to what CARA does.

#### Parameters:

- ppm\_f1: 1darray
  PPM scale of the indirect dimension
- t\_f1: 1darray
  Trace of the indirect dimension, real part
- SFO1: float
  Larmor frequency of the nucleus in the indirect dimension
- ppm\_f2: *1darray* PPM scale of the direct dimension
- t\_f2: 1darray
  Trace of the direct dimension, real part
- SFO2: float
  Larmor frequency of the nucleus in the direct dimension
- u\_1: float
  Chemical shift in F1 /ppm. Defaults to the center of the scale
- fwhm\_1: float Starting FWHM /Hz in the indirect dimension
- utol\_1: float
  Allowed tolerance for u 1 during the fit. (u 1-utol 1, u 1+utol 1)
- u\_2: float Chemical shift in F2 /ppm. Defaults to the center of the scale
- fwhm\_2: float Starting FWHM /Hz in the direct dimension
- utol\_2: float Allowed tolerance for u\_2 during the fit. (u\_2-utol\_2, u\_2+utol\_2)
- plot\_result: bool

  True to show how the program fitted the traces.

#### Returns:

• I\_tot: *float*Computed integral.

# 3.2.26 processing.integrate(fx, x=None, lims=None)

Calculates the definite integral of fx as I = F[-1] - F[0]. If fx is a multidimensional array, the integrals are computed along the last dimension.

#### Parameters:

- fx: ndarray
  Function (array) to integrate
- x: 1darray or None Independent variable. Determines the integration step. If None, it is the point scale
- lims: tuple or None Integration range. If None, the whole function is integrated.

#### Returns:

• I: float
Integrated function.

# 3.2.27 processing.interactive basl windows(ppm, data)

Allows for interactive partitioning of a spectrum in windows. Double left click to add a bar, double right click to remove it. Returns the location of the red bars as a list.

#### Parameters:

• ppm: 1darray

PPM scale of the spectrum

• data: 1darray

Spectrum to be partitioned

#### Returns:

• coord: *list*List containing the coordinates of the windows, plus ppm[0] and ppm[-1]

### 3.2.28 processing.interactive echo param(data0)

Interactive plot that allows to select the parameters needed to process a CPMG-like FID. Use the TextBox or the arrow keys to adjust the values. You can call processing.sum\_echo\_train or processing.split\_echo\_train by starring the return statement of this function, i.e.:

```
processing.sum_echo_train(data0, *interactive_echo_train(data0))
```

as they are in the correct order to be used in this way.

#### Parameters:

• data0: ndarray CPMG FID

- n: *int*Distance between one echo and the next one
- n\_echoes: *int* Number of echoes to sum/split
- i\_p: *int* Offset points from the start of the FID

## 3.2.29 processing.interactive fp(fid0, acqus, procs)

Perform the processing of a 1D NMR spectrum interactively. The GUI offers the opportunity to test different window functions, as well as different tdeff values and final sizes. The active parameters appear as blue text.

#### Parameters:

• fid0: *1darray* FID to process

• acqus: *dict*Dictionary of acquisition parameters

• procs: *dict*Dictionary of processing parameters

#### Returns:

• pdata: *1darray*Processed spectrum

• procs: *dict*Updated dictionary of processing parameters:

## 3.2.30 processing.interactive\_phase\_1D(ppmscale, S)

This function allow to adjust the phase of 1D spectra interactively. Use the mouse scroll to regulate the values.

#### Parameters:

- $\bullet$  ppm scale: 1darray ppm scale of the spectrum. Used to regulate the pivot position
- S: *1darray* Spectrum to be phased. Must be complex!

#### Returns:

• phased\_data: *1darray* Phased spectrum

## 3.2.31 processing.interactive phase 2D(ppm f1, ppm f2, S, hyper=True)

Interactively adjust the phases of a 2D spectrum S must be complex or hypercomplex, so BEFORE TO UNPACK

#### Parameters:

- ppm\_f1: *1darray* ppm scale of the indirect dimension
- ppm\_f2: 1darray ppm scale of the direct dimension
- S: 2darray
  Data to be phase-adjusted
- hyper: bool

  True if S is hypercomplex, False if S is just complex

- S: 2darray
  Phased data
- final\_values\_f1: tuple (p0\_f1, p1\_f1, pivot\_f1)
- final\_values\_f2: tuple (p0\_f2, p1\_f2, pivot\_f2)

## 3.2.32 processing.interactive\_qfil(ppm, data\_in)

Interactive function to design a gaussian filter with the aim of suppressing signals in the spectrum. You can adjust position and width of the filter scrolling with the mouse.

#### Parameters:

• ppm: *1darray* Scale on which the filter will be built

• data\_in: *1darray* Spectrum on which to apply the filter.

#### Returns:

• u: *float*Position of the gaussian filter

ullet s: float Width of the gaussian filter (Standard deviation)

## $3.2.33 \quad \text{processing.interactive} \\ \_\text{xfb} \\ (\text{fid0, acqus, procs, lvl0} \\ = 0.1, \\ \text{show} \\ \_\text{cnt} \\ = \\ \text{True})$

Perform the processing of a 2D NMR spectrum interactively. The GUI offers the opportunity to test different window functions, as well as different tdeff values and final sizes. The active parameters appear as blue text. When changing the parameters, give it some time to compute. The figure panel is quite heavy.

#### **Parameters:**

- fid0: 2darray
  FID to process
- acqus: *dict*Dictionary of acquisition parameters
- procs: *dict*Dictionary of processing parameters
- lvl0: *float* Starting level of the contours
- show\_cnt: bool Choose if to display data using contours (True) or heatmap (False)

- pdata: 2darray
  Processed spectrum
- procs: *dict*Updated dictionary of processing parameters

# 3.2.34 processing.inv\_convolve(in1, in2)

Perform the inverse-convolution of the two array by dividing their inverse Fourier transform. The two arrays must have the same dimension.

#### Parameters:

• in1: ndarray
First array

• in2: ndarray
Second array

#### Returns:

• cnv: ndarray
Convolved array

## 3.2.35 processing.inv fp(data, wf=None, size=None, fcor=0.5)

Performs the full inverse processing of a 1D NMR spectrum (data).

#### Parameters:

- data: *1darray* Spectrum
- wf: dict {'mode': function to be used, 'parameters': different from each function}
- ullet size: int initial size of the FID
- fcor: *float* weighting factor for the FID first point

#### Returns:

• pdata: *1darray* FID

# 3.2.36 processing.inv\_xfb(data, wf=[None, None], size=(None, None), fcor=[0.5, 0.5], FnMODE='States-TPPI')

Reverts the full processing of a 2D NMR FID (data).

#### Parameters:

• data: 2darray
Input data, hypercomplex

• wf: list of dict list of two entries [F1, F2]. Each entry is a dictionary of window functions

• size: *list of int*Initial size of FID

• fcor: *list of float* first fid point weighting factor [F1, F2]

• FnMODE: strAcquisition mode in F1

#### Returns:

• data: *2darray* Processed data

# 3.2.37 processing.iterCadzow(data, n, nc, itermax=100, f=0.005, print\_head=T print\_time=True)

This functions performs Cadzow denoising on  $\mathtt{data}$ , which is a 1D array of N points, in an iterative manner. The algorithm works as follows:

- 1. Transform data in a Hankel matrix  $\mathbb{H}$  of dimensions (N-n,n)
- 2. Make SVD on  $\mathbb{H} = \mathbb{USV}^{\dagger}$
- 3. Keep only the first nc singular values, and put all the rest to  $0 (S \to S')$
- 4. Rebuild  $\mathbb{H}' = \mathbb{U}\mathbb{S}'\mathbb{V}^{\dagger}$
- 5. Average the antidiagonals to rebuild the Hankel-type structure, then make 1D array
- 6. Check arrest criterion: if it is not reached, go to step 1, otherwise exit from the cycle and return the processed data.

The arrest criterion is on the array of singular values S, which is the main diagonal of the matrix S. At step k and Python indexing system:

$$\left|\frac{S^{(k-1)}[\text{nc}-1]}{S^{(k-1)}[\textbf{0}]} - \frac{S^{(k)}[\text{nc}-1]}{S^{(k)}[\textbf{0}]}\right| < f\frac{S^{(0)}[\text{nc}-1]}{S^{(0)}[\textbf{0}]}$$

#### **Parameters**

- data: *1darray* Input data
- n: *int*Number of columns of the Hankel matrix.
- nc: *int*Number of singular values to keep.
- itermax: *int*Maximum number of iterations allowed.
- f: float
  Factor for the arrest criterion.
- print\_head: *bool*Set it to True to display the fancy heading.
- print\_time: bool
  Set it to True to display the time spent.

#### Returns

• datap: *1darray* Denoised data

## 3.2.38 processing.load baseline(filename, ppm, data)

Read the baseline parameters from a file and builds the baseline itself.

#### Parameters:

 $\bullet$  filename: str Location of the baseline file

• ppm: *1darray* PPM scale of the spectrum

• data: *1darray*Spectrum of which to correct the baseline

#### Returns:

• baseline: *1darray* Computed baseline

## 3.2.39 processing.lp(data, pred=1, order=8, mode='b')

Apply linear prediction on the dataset. This method solves the linear system

$$\mathbb{D} \mathbf{a} = \mathbf{d}$$

where a is the array of lp coefficients.

#### Parameters:

• data: *1darray* 

FID to be linear-predicted

 $\bullet$  pred: int

Number of points to predict

 $\bullet$  order: int

Number of coefficients to use for the prediction

 $\bullet$  mode: str

'f' for forward linear prediction, 'b' for backward linear prediction

#### Returns:

• newdata: 1darray

FID with linear prediction applied.

## 3.2.40 processing.lrd(data, nc)

Denoising method based on Low-Rank Decomposition. The algorithm performs a singular value decomposition on data, then keeps only the first nc singular values while setting all the others to 0. Finally, rebuilds the data matrix using the modified singular values.

#### Parameters:

• data: *2darray*Data to be denoised

• nc: *int*Number of components, i.e. number of singular values to keep

#### Returns:

• data\_out: 2darray
Denoised data

## 3.2.41 processing.make polynomion baseline(ppm, data, limits)

Interactive baseline correction with 4th degree polynomion.

#### Parameters:

• ppm: 1darray

PPM scale of the spectrum

• data: *1darray* spectrum

 $\bullet$  limits: tuple

Window limits (left, right).

#### Returns:

 $\bullet$  mode: str

Baseline correction mode: 'polynomion' as default, 'spline' if you press the button

• C f: 1darray or str

Baseline polynomion coefficients, or 'callintsmooth' if you press the spline button

## 3.2.42 processing.make scale(size, dw, rev=True)

Computes the frequency scale of the NMR spectrum, given the # of points and the employed dwell time (the REAL one, not the TopSpin one!). 'rev'=True is required for the correct frequency arrangement in the NMR sense.

#### Parameters:

• size: *int*Number of points of the frequency scale

• dw : *float*Time spacing in the time dimension

• rev: bool
Reverses the scale

#### Returns:

• fqscale: *1darray*The computed frequency scale.

## 3.2.43 processing.mask sgn basl(ppm, data, SFO1, alpha=3, winsize=50)

Given an NMR spectrum, this function estimates the signal and baseline regions, and return a list of slices to cut the spectrum accordingly.

#### Parameters:

• ppm: *1darray* ppm scale of the spectrum

• data: *1darray* Spectrum

• SFO1: float Nucleus' Larmor frequency /MHz

• alpha: float
Factor that multiplies the std of the spectrum to set the threshold

• winsize: *float*Minimum size of the window that can contain peaks /Hz

#### **Returns:**

• peak\_slices: *list of slices*Slices that trim the data in the signal-only regions

• basl\_slices: *list of slices*Slices that trim the data in the baseline-only regions

# 3.2.44 processing.mcr(input\_data, nc, f=10, tol=1e-05, itermax=10000.0, P='H', oncols=True)

This is an implementation of Multivariate Curve Resolution for the denoising of 2D NMR data. Let us consider a matrix  $\mathbb{D}$ , of dimensions  $m \times n$ , where the starting data are stored. The final purpose of MCR is to decompose the  $\mathbb{D}$  matrix as follows:

$$\mathbb{D} = \mathbb{CS} + \mathbb{E}$$

where  $\mathbb{C}$  and  $\mathbb{S}$  are matrices of dimension  $m \times nc$  and  $nc \times n$ , respectively, and  $\mathbb{E}$  contains the part of the data that are not reproduced by the factorization. Being  $\mathbb{D}$  the FID of a NMR spectrum,  $\mathbb{C}$  will contain time evolutions of the indirect dimension, and  $\mathbb{S}$  will contain transients in the direct dimension.

The total MCR workflow can be separated in two parts: a first algorithm that produces an initial guess for the three matrices  $\mathbb{C}$ ,  $\mathbb{S}$  and  $\mathbb{E}$  (simplisma), and an optimization step that aims at the removal of the unwanted features of the data by iteratively filling the  $\mathbb{E}$  matrix (MCR ALS). This function returns the denoised datasets, CS, and the single C and S matrices.

#### Parameters:

- input\_data: 2darray or 3darray a 3D array containing the set of 2D NMR datasets to be coprocessed stacked along the first dimension. A single 2D array can be passed, if the denoising of a single dataset is desired.
- nc: *int* number of purest components to be looked for;
- f: float percentage of allowed noise;
- tol: *float* tolerance for the arrest criterion;
- itermax: *int* maximum number of allowed iterations
- P: str or 2darray
  'H' for horizontal stacking, 'V' for vertical stacking, or custom matrix as explained in the description of mcr stack
- oncols: bool

  True to estimate S with processing.simplisma, False to estimate C.

- CS\_f: 2darray or 3darray Final denoised data matrix
- C\_f: 2darray or 3darray Final C matrix
- S\_f: 2darray or 3darray Final S matrix

## 3.2.45 processing.mcr als(D, C, S, itermax=10000, tol=1e-05)

Performs alternating least squares to get the final  $\mathbb{C}$  and  $\mathbb{S}$  matrices. Being the fundamental MCR equation:  $\mathbb{D} = \mathrm{CS} + \mathrm{E}$  At the k-th step of the iterative cycle:

1. 
$$\mathbb{C}_k = \mathbb{DS}_{k-1}^+$$

2. 
$$\mathbb{S}_k = \mathbb{C}_k^+ \mathbb{D}$$

3. 
$$\mathbb{E}_k = \mathbb{D} - \mathbb{C}_k \mathbb{S}_k$$

Defined  $r_C$  and  $r_S$  as the Frobenius norm of the difference of C and S matrices between two subsequent steps:

$$rC = \|\mathbb{C}_k - \mathbb{C}_{k-1}\| \quad rS = \|\mathbb{S}_k - \mathbb{S}_{k-1}\|$$

The convergence is reached when: rC <= tol && rS <= tol

#### Parameters:

- D: *2darray*Input data, of dimensions m x n
- C: 2darray Estimation of the C matrix, of dimensions m x nc.
- S: 2darray
  Estimation of the S matrix, of dimensions nc x n.
- itermax: *int*Maximum number of iterations
- tol: *float*Threshold for the arrest criterion.

- C: 2darray
  Optimized C matrix, of dimensions m x nc.
- S: 2darray
  Optimized S matrix, of dimensions nc x n.

### 3.2.46 processing.mcr stack(input data, P='H')

Performs matrix augmentation by assembling input\_data according to the positioning matrix P. P has two default modes: 'H' = horizontal stacking; 'V' = vertical stacking. Otherwise, a custom  $\mathbb P$  matrix can be given as follows. The entries of the  $\mathbb P$  matrix are the indices of the data in input\_data. The shape of the matrix determines the final arrangement.

Example: if input\_data is [a, b, c, d, e, f], and one wants to obtain [[a, b], [d,c], [f, e]] the correspondant P matrix is:

$$P = [[0, 1], [3, 2], [5, 4]]$$

If each dataset in input\_data has dimensions (m, n) and P has dimensions (u,v), then the returned data matrix will have dimensions (mu, nv).

#### **Parameters:**

- input\_data: 3darray
  Contains the spectra to be stacked together. The index that runs on the datasets must be the first one.
- P: str or 2darray
  'H' for horizontal stacking, 'V' for vertical stacking, or custom matrix as explained in the description

#### Returns:

• data: *2darray*Augmented data matrix.

## 3.2.47 processing.mcr unpack(C, S, nds, P='H')

Reverts matrix augmentation of mcr\_stack. The denoised spectra can be calculated by matrix multiplication:  $D[k] = C_f[k] S_f[k] \# for k = 0, ..., nds-1$ 

#### Parameters:

- C: 2darray MCR C matrix
- S: 2darray MCR S matrix
- nds: *int* number of experiments
- P: str or 2darray
  'H' for horizontal stacking, 'V' for vertical stacking, or custom matrix as explained in the description of mcr stack

- C\_f: list of 2darray Disassembled MCR C matrix
- S\_f: *list of 2darray* Disassembled MCR C matrix

# $3.2.48 \quad processing.pknl(data, grpdly=0, onfid=False)$

Compensate for the Bruker group delay at the beginning of FID through a first-order phase correction of p1 = 360 \* GRPDLY This should be applied after apodization and zero-filling.

#### Parameters:

• data: *ndarray*Input data. Be sure it is complex!

• grpdly: *int*Number of points that make the group delay.

• on fid: bool If it is True, performs FT before to apply the phase correction, and IFT after.

#### Returns:

• datap: ndarray Corrected data

# 3.2.49 processing.ps(data, ppmscale=None, p0=None, p1=None, pivot=None, interactive=False)

Applies phase correction on the last dimension of data. The pivot is set at the center of the spectrum by default. Missing parameters will be inserted interactively.

#### Parameters:

• data: ndarray Input data

• ppmscale: 1darray or None PPM scale of the spectrum. Required for pivot and interactive phase correction

• p0: float
Zero-order phase correction angle /degrees

• p1: float
First-order phase correction angle /degrees

• pivot: float or None. First-order phase correction pivot /ppm. If None, it is the center of the spectrum.

• interactive: bool

If True, all the parameters will be ignored and the interactive phase correction panel will be opened.

#### Returns:

• datap: ndarray Phased data

• final\_values: tuple Employed values of the phase correction. (p0, p1, pivot)

## 3.2.50 processing.qfil(ppm, data, u, s)

Suppress signals in the spectrum using a gaussian filter.

#### Parameters:

• ppm: 1darray

Scale on which to build the filter

• data: ndarray

Data to be processed. The filter is applied on the last dimension

• u: float

Position of the filter

• s: float

Width of the filter (standard deviation)

#### Returns:

• pdata: ndarray Filtered data

# 3.2.51 processing.qpol(fid)

Fits the FID with a 4-th degree polynomion, then subtracts it from the original FID. The real and imaginary channels are treated separately.

#### Parameters:

• fid : ndarray Self-explanatory.

#### Returns:

• fid\_corr : ndarray Processed FID

# $3.2.52 \quad \text{processing.qsin(data, ssb)}$

Sine-squared apodization.

### Parameters:

• ssb: *int*Sine bell shift.

# 3.2.53 processing.quad(fid)

Subtracts from the FID the arithmetic mean of its last quarter. The real and imaginary channels are treated separately.

#### Parameters:

• fid : ndarray Self-explanatory.

#### Returns:

• fid : ndarray Processed FID.

## 3.2.54 processing.repack 2D(rr, ir, ri, ii)

Renconstruct hypercomplex 2D NMR data given the 4 ser files

#### Parameters:

- rr: 2darray Real F2, Real F1
- ir: 2darray Imaginary F2, Real F1
- ri: 2darray Real F2, Imaginary F1
- ii: 2darray Imaginary F2, Imaginary F1

#### Returns:

• data: *2darray* Hypecomplex matrix

# 3.2.55 processing.rev(data)

Reverse data over its last dimension

# 3.2.56 processing.rpbc(data, split\_imag=False, n=5, basl\_method='huber', basl\_thresh=0.2, basl\_itermax=2000, \*\*phase\_kws)

Reversed Phase and Baseline Correction. Allows for the automatic phase correction and baseline subtraction of NMR spectra. It is called 'reversed' because the baseline is actually computed and subtracted before to perform the phase correction.

The baseline is computed using a low-order polynomion, built on a scale that goes from -1 to 1, whose coefficients are obtained minimizing a non-quadratic cost function. It is recommended to use either 'tq' (truncated quadratic, much faster) or 'huber' (Huber function, slower but sometimes more accurate). The user is requested to choose between separating the real and imaginary channel in this step. The order of the polynomion and the threshold value are the key parameters for obtaining a good baseline. The used function is processing.polyn\_basl

The phase correction is computed on the baseline-subtracted complex data as described in the SINC algorithm (ref.). The default parameters are generally fine, but in case of data with poor SNR (approximately SNR < 10) better results can be obtained by increasing the value of the e1 parameter. The employed function is processing.SINC\_phase

#### Parameters:

- data: *1darray*Data to be processed, complex-valued
- split\_imag: bool

  If True, computes the baseline on the real and imaginary part separately; else, the set of polynomion coefficients are forced to be the same for both
- $\bullet$  n: intNumber of coefficients of the polynomion, i.e. it will be of degree n-1
- basl\_method: str Cost function to be minimized for the baseline computation. Look for fit.CostFunc, 'method' attribute
- basl\_thresh: float
  Relative threshold value for the non-quadratic behaviour of the cost function. Look for fit.CostFunc,
  's' attribute
- basl\_itermax: int

  Maximum number of iterations allowed during the baseline fitting procedure
- phase\_kws: keyworded arguments
  Optional arguments for the phase correction. Look for fit.SINC\_phase keyworded arguments
  for details.

- y: *1darray* Processed data
- p0: float
  Zero-order phase correction angle, in degrees
- p1: float
  First-order phase correction angle, in degrees

• c: 1darray

Set of coefficients to be used for the baseline computation, starting from the 0-order coefficient

### 3.2.57 processing.simplisma(D, nc, f=10, oncols=True)

Finds the first nc purest components of matrix  $\mathbb D$  using the simplisma algorithm, proposed by Windig and Guilment (DOI: 10.1021/ac00014a016). If oncols=True, this function estimates  $\mathbb S$  with simplisma, then calculates  $\mathbb C = \mathbb D\mathbb S^+$ . If oncols=False, this function estimates  $\mathbb C$  with simplisma, then calculates  $\mathbb S = \mathbb C^+\mathbb D$ . If defines the percentage of allowed noise.

#### **Parameters:**

- D: 2darray
  Input data, of dimensions m x n
- nc: int
  Number of components to be found. This determines the final size of the C and S matrices.
- f: float
  Percentage of allowed noise.
- oncols: bool

  If True, simplisma estimates the S matrix, otherwise estimates C.

- C: 2darray Estimation of the C matrix, of dimensions m x nc.
- S: 2darray
  Estimation of the S matrix, of dimensions nc x n.

# 3.2.58 processing. $\sin(\text{data, ssb})$

Sine apodization.

### Parameters:

• ssb: *int*Sine bell shift.

## 3.2.59 processing.smooth\_g(d, m)

Apply a smoothing with a gaussian filter by convolution. The width of the filter is 1/m.

#### Parameters:

- d: *1darray*Data to be smoothed
- m: *float*Inverse width of the filter /pt

#### Returns:

• yc: *1darray* Smoothed data

## 3.2.60 processing.split echo train(datao, n, n echoes, i p=0)

Separate a CPMG echo-train FID into echoes so to be processed separately. The first decay, i.e. the native FID, is extracted, and corresponds to echo number 0. Then, for each echo, the left side (reversed) is summed up to its right part.

#### Parameters:

- datao: *ndarray* FID with an echo train on its last dimension
- n: *int* number of points that separate one echo from the next
- n\_echoes: *int* number of echoes to extract. If it is 0, extracts only the first decay
- i\_p: *int*Number of offset points

#### Returns:

• data\_p: (n+1)darraySeparated echoes

## 3.2.61 processing.stack fids(\*fids, filename=None)

Stacks together FIDs in order to create a pseudo-2D experiment. This function can handle either arrays or Spectrum\_1D objects.

#### Parameters:

- fids: sequence of 1darrays or Spectrum\_1D objects Input data.
- $\bullet$  filename: str Location for a .npy file to be saved. If None, no file is created.

#### Returns:

• p2d: 2darray Stacked FIDs.

## 3.2.62 processing.sum echo train(datao, n, n echoes, i p=0)

Sum up a CPMG echo-train FID into echoes so to be enchance the SNR. This function calls processing.split\_echo\_train with the same parameters.

#### Parameters:

- datao: *ndarray* FID with an echo train on its last dimension
- $\bullet$  n: int number of points that separate one echo from the next
- n\_echoes: *int* number of echoes to sum
- i\_p: *int*Number of offset points

#### Returns:

• data\_p: ndarray Summed echoes

# 3.2.63 processing.td\_eff(data, tdeff)

Uses only the first tdeff points of data. tdeff must be a list as long as the dimensions: tdeff = [F1, F2, ..., Fn]

#### Parameters:

• data: *ndarray*Data to be trimmed

• tdeff: *list of int*Number of points to be used in each dimension

# 3.2.64 processing.tp\_hyper(data)

Computes the hypercomplex transpose of data. Needed for the processing of data acquired in a phase-sensitive manner in the indirect dimension.

# 3.2.65 processing.unpack\_2D(data)

Separates hypercomplex data into 4 distinct ser files

#### Parameters:

• data: *2darray* Hypercomplex matrix

#### Returns:

- rr: 2darray Real F2, Real F1
- ir: 2darray Imaginary F2, Real F1
- ri: *2darray* Real F2, Imaginary F1
- ii: 2darray Imaginary F2, Imaginary F1

## 3.2.66 processing.whittaker smoother(data, n=2, s f=1, w=None)

Adapted from P.H.C. Eilers, Anal. Chem 2003, 75, 3631-3636. Implementation of the smoothing algorithm proposed by Whittaker in 1923.

#### Parameters:

- data: *1darray*Data to be smoothed
- $\bullet$  n: int Order of the difference to be computed
- s\_f: *float* Smoothing factor
- w: 1darray or None Array of weights. If None, no weighting is applied.

#### Returns:

• z: 1darray Smoothed data

## 3.2.67 processing.write basl info(f, limits, mode, data)

Writes the baseline parameters of a certain window in a file.

- f: TextIO object File where to write the parameters
- limits: *tuple*Limits of the spectral window. (left, right)
- mode: str Baseline correction mode: 'polynomion' or 'spline'
- data: float or 1darray
  It can be either the spline smoothing factor or the polynomion coefficients

# 3.2.68 processing.xfb(data, wf=[None, None], zf=[None, None], fcor=[0.5, 0.5], tdeff=[0, 0], u=True, FnMODE='States-TPPI')

Performs the full processing of a 2D NMR FID (data). The returned values depend on u: it is True, returns a sequence of 2darrays depending on FnMODE, otherwise just the complex/hypercomplex data after FT in both dimensions

#### **Parameters:**

- data: *2darray* Input data
- wf: sequence of dict (F1, F2); {'mode': function to be used, 'parameters': different from each function}
- zf: sequence of int final size of spectrum, (F1, F2)
- fcor: sequence of float weighting factor for the FID first point, (F1, F2)
- tdeff: sequence of int number of points of the FID to be used for the processing, (F1, F2)
- u: bool choose if to unpack the hypercomplex spectrum into separate arrays or not
- FnMODE: strAcquisition mode in F1

#### Returns:

• datap: 2darray or tuple of 2darray Processed data or tuple of 2darray

# 3.2.69 processing.zf(data, size)

Zero-filling of data up to size in its last dimension.

#### Parameters:

• data: ndarrayArray to be zero-filled

ullet size: int Number of points of the last dimension after zero-filling

#### Returns:

• datazf: ndarray Zero-filled data

# 3.3 FIGURES package

This package contains a series of functions to make plots of various nature.

3.3.1 figures.ax1D(ax, ppm, datax, norm=False, xlims=None, ylims=None, c='tab:blue', lw=0.5, X\_label=, Y\_label='Intensity /a.u.', n\_xticks=10, n\_yticks=10, label=None, fontsize=10)

Makes the figure of a 1D NMR spectrum, placing it in a given figure panel. This allows the making of modular figures.

The plot can be customized in a very flexible manner by setting the function keywords properly.

- ax: matplotlib.subplot Object panel where to put the figure
- ppm: *1darray* ppm scale of the spectrum
- data: *1darray* spectrum to be plotted
- norm: bool if True, normalizes the intensity to 1.
- xlims: *list or tuple*Limits for the x-axis. If None, the whole scale is used.
- ylims: *list or tuple*Limits for the y-axis. If None, the whole scale is used.
- c: str Colour of the line.
- lw: float linewidth
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int* Number of numbered ticks on the x-axis of the figure
- label: str label to be put in the legend.
- fontsize: *float*Biggest font size in the figure.

### Returns:

 $\bullet$  line: Line2D Object Line object returned by plt.plot.

 $\begin{array}{lll} 3.3.2 & \text{figures.ax2D(ax, ppm\_f2, ppm\_f1, datax, xlims=None, ylims=None,} \\ & \text{cmap='Greys\_r', c\_fac=1.4, lvl=0.1, lw=0.5, X\_label=, Y\_label=,} \\ & \text{title=None, n xticks=10, n yticks=10, fontsize=10)} \end{array}$ 

Makes a 2D contour plot like the one in figures.figure2D, but in a specified panel. Allows for the buildup of modular figures. The contours are drawn according to the formula:

cl = contour\\_start \* contour\\_factor \*\* np.arange(contour\\_num)

where contour\_start = np.max(data)\* lvl, contour\_num = 16 and contour\_factor = c\\_fac. Increasing the value of c\_fac will decrease the number of contour lines, whereas decreasing the value of c fac will increase the number of contour lines.

- ax: matplotlib.subplot Object panel where to put the figure
- ppm\_f2: *1darray* ppm scale of the direct dimension
- ppm\_f1: *1darray* ppm scale of the indirect dimension
- datax: 2darray the 2D NMR spectrum to be plotted
- xlims: *tuple* limits for the x-axis (left, right). If None, the whole scale is used.
- ylims: tuple limits for the y-axis (left, right). If None, the whole scale is used.
- cmap: strColormap identifier for the contour
- c\_fac: *float* Contour factor parameter
- lvl: *float* height with respect to maximum at which the contour are computed
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- lw: *float* linewidth of the contours
- title: str Figure title.
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure

- $\bullet\,$  n\_yticks: int Number of numbered ticks on the x-axis of the figure
- fontsize: *float*Biggest font size in the figure.

#### Returns:

• cnt: matplotlib.QuadContour object Drawn contour lines

3.3.3 figures.ax\_heatmap(ax, data, zlim='auto', z\_sym=True, cmap=None, xscale=None, yscale=None, rev=(False, False), n\_xticks=10, n\_yticks=1 n zticks=10, fontsize=10)

Computes a heatmap of data on the given 'ax'

#### Parameters:

- ax: matplotlib.Subplot object
  Panel where to draw the heatmap
- data: *2darray* Input data
- zlim: tuple or 'auto' or 'abs'
  Vertical limits of the heatmap, that determines the extent of the colorbar. 'auto' means (min(data), max(data)), 'abs' means(min(|data|), max(|data|)).
- z\_sym: bool

  True to symmetrize the vertical scale around 0.
- cmap: matplotlib.cm object Colormap of the heatmap.
- xscale: 1darray or None x-scale. None means np.arange(data.shape[1])
- yscale: *1darray or None* y-scale. None means np.arange(data.shape[0])
- rev: tuple of bool Reverse scale (x, y).
- n\_xticks: *int*Number of ticks of the x axis
- n\_yticks: *int*Number of ticks of the y axis
- n\_zticks: *int*Number of ticks of the color bar
- fontsize: *float*Biggest font size to apply to the figure.

#### **Returns:**

- im: matplotlib.AxesImageThe heatmap
- cax: figure panel where the colorbar is drawn

## 3.3.4 figures.dotmd(ppmscale, S, labels=None, lw=0.8, n xticks=10)

Interactive display of multiple 1D spectra.

#### Parameters:

- ppmscale: 1darray or list ppm scale of the spectra. If only one scale is supplied, all the spectra are plotted using the same scale. Otherwise, each spectrum is plotted using its scale.
- S: *list* spectra to be plotted
- labels: *list* labels to be put in the legend.
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure

#### Returns:

• scale\_factor: *list*Intensity of the spectra with respect to the original when the figure is closed

3.3.5 figures.dotmd\_2D(ppm\_f1, ppm\_f2, S0, labels=None, name='dotmd\_2I X\_label='\$ delta \$ F2 /ppm', Y\_label='\$ delta \$ F1 /ppm', n xticks=10, n yticks=10, Neg=False)

Interactive display of multiple 2D spectra. They have to share the same scales.

#### **Parameters:**

- ppm\_f1: 1darray ppm scale of the indirect dimension. If only one scale is supplied, all the spectra are plotted using the same scale. Otherwise, each spectrum is plotted using its scale. There is a 1:1 correspondence between ppm f1 and S.
- ppm\_f2: 1darray ppm scale of the direct dimension. If only one scale is supplied, all the spectra are plotted using the same scale. Otherwise, each spectrum is plotted using its scale. There is a 1:1 correspondence between ppm f2 and S.
- S: *list* spectra to be plotted
- labels: *list* labels to be put in the legend.
- name: *str*If you choose to save the figure, this is its filename.
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- n\_xticks: *int*Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int* Number of numbered ticks on the x-axis of the figure
- Neg: bool

  If True, show the negative contours.

#### Returns:

• lvl: *list*Intensity factors when the figure is closed

3.3.6 figures.figure1D(ppm, datax, norm=False, xlims=None, ylims=None, c='tab:blue', lw=0.5,  $X_label=$ ,  $Y_label=$ 'Intensity /a.u.', n\_xticks=10, n\_yticks=10, fontsize=10, name=None, ext='tiff', dpi=600)

Makes the figure of a 1D NMR spectrum.

The plot can be customized in a very flexible manner by setting the function keywords properly.

- ppm: *1darray* ppm scale of the spectrum
- datax: *1darray* spectrum to be plotted
- norm: bool if True, normalizes the intensity to 1.
- xlims: *list or tuple*Limits for the x-axis. If None, the whole scale is used.
- ylims: *list or tuple*Limits for the y-axis. If None, the whole scale is used.
- c: str Colour of the line.
- lw: float linewidth
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- n\_xticks: *int*Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int* Number of numbered ticks on the x-axis of the figure
- fontsize: *float*Biggest font size in the figure.
- name:  $str\ or\ None$ Filename for the figure to be saved. If None, the figure is shown instead.
- ext: strFormat of the image
- dpi: *int*Resolution of the image in dots per inches

3.3.7 figures.figure1D\_multi(ppm0, data0, xlims=None, ylims=None, norm=Fac=None, X\_label=, Y\_label='Intensity /a.u.', n\_xticks=10, n\_yticks=1 fontsize=10, labels=None, name=None, ext='tiff', dpi=600)

Creates the superimposed plot of a series of 1D NMR spectra.

- ppm0: sequence of 1darray or 1darray ppm scale of the spectra. If only one scale is supplied, it is assumed to be the same for all the spectra
- data0: sequence of 1darray
  List containing the spectra to be plotted
- xlims: tuple or None
  Limits for the x-axis. If None, the whole scale is used.
- ylims: tuple or None
  Limits for the y-axis. If None, they are automatically set.
- norm: False or float or str

  If it is False, it does nothing. If it is float, divides all spectra for that number. If it is str('#'), normalizes all the spectra to the '#' spectrum (python numbering). If it is whatever else string, normalizes all spectra to themselves.
- c: tuple or None
  List of the colors to use for the traces. None uses the default ones.
- X\_label: str text of the x-axis label
- Y\_label: str text of the y-axis label
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int*Number of numbered ticks on the x-axis of the figure
- fontsize: *float*Biggest fontsize in the picture
- labels: list or None or False
  List of the labels to be shown in the legend. If it is None, the default entries are used (i.e., '1, 2, 3,...'). If it is False, the legend is not shown.
- name:  $str\ or\ None$  Filename of the figure, if it has to be saved. If it is None, the figure is shown instead.
- ullet ext: str Format of the image
- dpi: *int*Resolution of the image in dots per inches

3.3.8 figures.figure2D(ppm\_f2, ppm\_f1, datax, xlims=None, ylims=None, cmap='Greys\_r', c\_fac=1.4, lvl=0.09, X\_label=, Y\_label=, lw=0.5, cmapneg=None, n\_xticks=10, n\_yticks=10, fontsize=10, name=None, ext='tiff', dpi=600)

Makes a 2D contour plot. Allows for the buildup of modular figures. The contours are drawn according to the formula:  $cl = contour\_start * contour\_factor ** np.arange(contour\_num)$  where contour\\_start = np.max(data) \* lvl, contour\\_num = 16 and contour\\_factor = c\_fac. Increasing the value of c\_fac will decrease the number of contour lines, whereas decreasing the value of c\_fac will increase the number of contour lines.

- ppm\_f2: *1darray* ppm scale of the direct dimension
- ppm\_f1: *1darray* ppm scale of the indirect dimension
- datax: 2darray the 2D NMR spectrum to be plotted
- xlims: tuple limits for the x-axis (left, right). If None, the whole scale is used.
- ylims: tuple limits for the y-axis (left, right). If None, the whole scale is used.
- cmap: strColormap identifier for the contour
- c\_fac: *float* Contour factor parameter
- lvl: *float* height with respect to maximum at which the contour are computed
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- lw: *float* linewidth of the contours
- cmapneg: str or None Colormap identifier for the negative contour. If None, they are not computed at all
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int* Number of numbered ticks on the x-axis of the figure

ullet fontsize: float

Biggest font size in the figure.

 $\bullet$  name: str

Filename for the figure

 $\bullet$  ext: str

Format of the image

 $\bullet$  dpi: int

Resolution of the image in dots per inches

 $\begin{array}{lll} 3.3.9 & \text{figures.figure2D\_multi(ppm\_f2, ppm\_f1, datax, xlims=None, ylims=None, lvl='default', c\_fac=1.4, Negatives=False, X\_label=, Y\_label=, lw=0.5, n\_xticks=10, n\_yticks=10, labels=None, name=None, ext='tiff' dpi=600) \end{array}$ 

Generates the figure of multiple, superimposed spectra, using figures.ax2D.

- ppm\_f2: 1darray ppm scale of the direct dimension
- ppm\_f1: *1darray* ppm scale of the indirect dimension
- datax: sequence of 2darray the 2D NMR spectra to be plotted
- xlims: *tuple* limits for the x-axis (left, right). If None, the whole scale is used.
- ylims: *tuple* limits for the y-axis (left, right). If None, the whole scale is used.
- lvl: 'default' or list height with respect to maximum at which the contour are computed. If 'default', each spectrum is at 10
- c\_fac: *float*Contour factor
- Negatives: bool set it to True if you want to see the negative part of the spectrum
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- lw: *float* linewidth of the contours
- n\_xticks: *int*Number of numbered ticks on the x-axis of the figure
- n\_yticks: *int* Number of numbered ticks on the x-axis of the figure
- labels: *list* entries of the legend. If None, the spectra are numbered.
- name: strFilename for the figure. If None, it is shown instead of saved
- ext: strFormat of the image

ullet dpi: int Resolution of the image in dots per inches

# $\begin{array}{ll} 3.3.10 & \text{figures.fitfigure}(S, ppm\_scale, t\_AQ, V, C=False, SFO1=701.125, \\ & o1p=0, \ limits=None, \ s\_labels=None, \ X\_label=, \ n\_xticks=10, \\ & name=None) \end{array}$

Makes the figure to show the result of a quantitative fit.

- S: *1darray* Spectrum to be fitted
- ppm\_scale : 1darray Self-explanatory
- V : 2darray matrix (# signals, parameters)
- C: 1darray or False

  Coefficients of the polynomion to be used as baseline correction. If the 'baseline' checkbox in the interactive figure panel is not checked, C f is False.
- limits: tuple or None
  Trim limits for the spectrum (left, right). If None, the whole spectrum is used.
- s\_labels: list or None or False Legend entries for the single components. If None, they are computed automatically as 1, 2, 3, etc. If False, they are not shown in the legend.
- X\_label : str label for the x-axis.
- n\_xticks: int number of numbered ticks that will appear in the ppm scale. An oculated choice can be very satisfying.
- name : str or None Name with which to save the figure. If None, the picture is shown instead of being saved.

3.3.11 figures.heatmap(data, zlim='auto', z\_sym=True, cmap=None, xs-cale=None, yscale=None, rev=(False, False), n\_xticks=10, n\_yticks=10 n\_zticks=10, fontsize=10, name=None)

Computes a heatmap of data.

- data: 2darray Input data
- zlim: tuple or 'auto' or 'abs'

  Vertical limits of the heatmap, that determines the extent of the colorbar. 'auto' means (min(data), max(data)), 'abs' means(min(|data|), max(|data|)).
- z\_sym: bool

  True to symmetrize the vertical scale around 0.
- cmap: matplotlib.cm object Colormap of the heatmap.
- xscale: *1darray or None* x-scale. None means np.arange(data.shape[1])
- yscale: *1darray or None* y-scale. None means np.arange(data.shape[0])
- rev: tuple of bool Reverse scale (x, y).
- n\_xticks: *int* Number of ticks of the x axis
- n\_yticks: *int* Number of ticks of the y axis
- n\_zticks: *int*Number of ticks of the color bar
- fontsize: *float*Biggest font size to apply to the figure.
- name: *str or None*Filename for the figure. Set to None to show the figure.

# $\begin{array}{ll} \textbf{3.3.12} & \textbf{figures.ongoing\_fit(exp, calc, residual, ylims=None, filename=None,} \\ & \textbf{dpi}{=}100) \end{array}$

Makes a figure of an ongoing fit. It displays the experimental data and the model, and the residuals in a separate window. The figure can be either saved or shown.

#### Parameters:

• exp: *1darray* Experimental data

• calc: *1darray*Current model

• residual: *1darray* Residuals of the fit

• ylims: *tuple* Optional limits for y-axis

• filename:  $str\ or\ None$ Filename of the figure to be saved. If None, the figure is shown instead

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

# 3.3.13 figures.plot\_fid(fid, name=None, ext='tiff', dpi=600)

Makes a two-panel figure that shows on the left the real part of the FID, on the right the imaginary part. The x-scale and y-scale are automatically adjusted.

# 3.3.14 figures.plot\_fid\_re(fid, scale=None, c='b', lims=None, name=None, ext='tiff', dpi=600)

Makes a single-panel figure that shows either the real or the imaginary part of the FID. The x-scale and y-scale are automatically adjusted.

- fid: ndarray
  FID to be plotted
- scale: *1darray or None* x-scale of the figure
- $\bullet$  c: str Color
- lims: tuple or None Limits
- name: strName of the figure
- ext: strFormat of the image
- dpi: *int*Resolution of the image in dots per inches

# 3.3.15 figures.redraw\_contours(ax, ppm\_f2, ppm\_f1, S, lvl, cnt, Neg=False, Ncnt=None, lw=0.5, cmap=[None, None])

Redraws the contours in interactive 2D visualizations.

#### Parameters:

- ax: matplotlib.Subplot Object
  Panel of the figure where to draw the contours
- ppm\_f2: 1darray ppm scale of the direct dimension
- ppm\_f1: *1darray* ppm scale of the indirect dimension
- S: 2darray Spectrum
- lvl: *float*Level at which to draw the contours
- cnt: matplotlib.contour.QuadContourSet object Pre-existing contours
- Neg: bool
  Choose if to draw the negative contours (True) or not (False)
- Ncnt: matplotlib.contour.QuadContourSet object Pre-existing negative contours
- lw: float Linewidth
- cmap: *list*Colour of the contours. [cmap +, cmap -]

#### Returns:

- cnt: matplotlib.contour.QuadContourSet object Updated contours
- Ncnt: matplotlib.contour.QuadContourSet object or None Updated negative contours if Neg is True, None otherwise

### 3.3.16 figures.sns heatmap(data, name=None, ext='tiff', dpi=600)

Computes a heatmap of data, which is a matrix. This function employs the seaborn package. Specify name if you want to save the figure.

#### Parameters:

• data: 2darray

Data of which to compute the heatmap. Make sure the entries are real numbers.

• name:  $str\ or\ None$ Filename of the figure to be saved. If None, the figure is shown instead.

• ext: strFormat of the image

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

 $\begin{array}{lll} 3.3.17 & \text{figures.stacked\_plot(ppmscale, S, xlims=None, lw=0.5, X\_label=,} \\ & & Y\_label='Normalized intensity / a.u.', n\_xticks=10, labels=None, \\ & & name=None, ext='tiff', dpi=600) \end{array}$ 

Creates a stacked plot of all the spectra contained in the list S. Note that S MUST BE a list. All the spectra must share the same scale.

- ppmscale: *1darray* ppm scale of the spectrum
- S: *list* spectra to be plotted
- xlims: *list or tuple*Limits for the x-axis. If None, the whole scale is used.
- lw: float linewidth
- name: *str* filename of the figure, if it has to be saved;
- X\_label: str text of the x-axis label;
- Y\_label: str text of the y-axis label;
- n\_xticks: *int* Number of numbered ticks on the x-axis of the figure
- labels: *list* labels to be put in the legend.

# 3.4 SIM package

This package contains function for the simulation of various features of NMR spectra, being them monodimensional or bidimensional. Functions for the simulation of whole spectra are also provided.

# 3.4.1 sim.calc splitting(u0, I0, m=1, J=0)

Calculate the frequency and the intensities of a NMR signal splitted by scalar coupling.

#### Parameters:

- u0: float
  Frequency of the non-splitted signal (Hz)
- I0: float
  Total intensity of the non-splitted signal.
- m: int
   Multiplicity, i.e. number of expected signals after the splitting
- J: float
  Scalar coupling constant (Hz)

#### Returns:

- u\_s: 1darray Frequencies of the splitted signal (Hz)
- I\_s: *1darray* Intensities of the splitted signal

3.4.2	sim.cron	func,	*args,	**kwargs)

Decorator: use it to monitor the runtime of a function.

# $3.4.3 \quad sim.f_gaussian(x, u, s, A=1)$

Gaussian function in the frequency domain:

#### Parameters:

- x: 1darray Independent variable
- ullet u: float Peak position
- s: *float* Standard deviation
- A: float Intensity

#### Returns:

• f: *1darray* Gaussian function.

# $3.4.4 \quad sim.f_lorentzian(x, u, fwhm, A=1)$

Lorentzian function in the time domain:

#### Parameters:

- x: 1darray Independent variable
- u: float Peak position
- ullet fwhm: float Full-width at half-maximum
- A: float Intensity

#### Returns:

• f: *1darray* Lorentzian function.

# $3.4.5 \quad sim.f_pvoigt(x, u, fwhm, A=1, b=0)$

Pseudo-Voigt function in the frequency domain:

#### Parameters:

- x: 1darray Independent variable
- u: float Peak position
- $\bullet$  fwhm: float Full-width at half-maximum
- A: float Intensity
- b: *float*Fraction of gaussianity

#### Returns:

• S: *1darray* Pseudo-Voigt function.

# $3.4.6 \quad sim.gaussian\_filter(ppm, u, s)$

Compute a gaussian filter to be used in order to suppress signals in the spectrum.

#### Parameters:

- ppm: *1darray* Scale on which to build the filter
- u: *float*Position of the filter
- s: *float*Width of the filter (standard deviation)

#### Returns:

• G: *1darray* Computed gaussian filter

# $3.4.7 \quad sim.load\_sim\_1D(File)$

Creates a dictionary from the spectral parameters listed in the input file.

#### Parameters:

 File: str Path to the input file location

#### Returns:

• dic: dict Dictionary of the parameters, ready to be read from the simulation functions.

## 3.4.8 sim.load sim 2D(File, states=True)

Creates a dictionary from the spectral parameters listed in the input file.

#### Parameters:

ullet File: str Path to the input file location

• states: bool

If FnMODE is States or States-TPPI, set it to True to get the correct timescale.

#### **Returns:**

• dic: dict
Dictionary of the parameters, ready to be read from the simulation functions.

 $3.4.9 \quad sim.mult\_noise(data\_size, mean, s\_n)$ 

Multiplicative noise model.

## 3.4.10 sim.multiplet(u, I, m='s', J=[])

Split a given signal according to a scalar coupling pattern.

#### Parameters:

- ullet u: float Frequency of the non-splitted signal (Hz)
- I: *float*Intensity of the non-splitted signal
- $\bullet\,$ m: str Organic chemistry-like multiplet, i.e. s, d, dqt, etc.
- J: float or list
  Scalar coupling constants. The number of constants should match the number of coupling branches

#### Returns:

- u\_in: *list*List of the splitted frequencies (Hz)
- I\_in: *list*Intensities of the splitted signal

# 3.4.11 sim.noisegen(size, o2, t2, s\_n=1)

Simulates additive noise in the time domain.

#### Parameters:

- size: int or tuple
  Dimension of the noise matrix
- o2: float Carrier frequency, in Hz.
- t2: *1darray*Time scale of the last temporal dimension.
- s\_n: *float* Standard deviation of the noise.

#### Returns:

• noise: 2darray
Noise matrix, of dimensions size.

# $3.4.12 \quad sim.sim\_1D(File, pv=False)$

Simulates a 1D NMR spectrum from the instructions written in File.

#### Parameters:

- File: str Path to the input file location
- pv: bool True for pseudo-Voigt model, False for Voigt model.

#### Returns:

• fid: *1darray* FID of the simulated spectrum.

## 3.4.13 sim.sim 2D(File, states=True, alt=True, pv=False)

Simulates a 2D NMR spectrum from the instructions written in File. The indirect dimension is sampled with states-TPPI as default.

#### Parameters:

• File: strPath to the input file location

• states: bool Set it to True to allow for correct spectral arrangement in the indirect dimension.

• alt: bool
Set it to True to allow for correct spectral arrangement in the indirect dimension.

• pv: bool
True for pseudo-Voigt model, False for Voigt model.

#### Returns:

• fid: 2darray FID of the simulated spectrum.

## 3.4.14 sim.t 2Dgaussian(t1, t2, v1, v2, s1, s2, A=1, states=True, alt=True)

Bidimensional gaussian function.

#### Parameters:

- t1: *1darray*Indirect evolution timescale
- t2: *1darray*Timescale of the direct dimension
- v1: float
  Peak position in the indirect dimension, in Hz
- v2: float
  Peak position in the direct dimension, in Hz
- s1: *float* Standard deviation in the indirect dimension, in rad/s
- s2: float Standard deviation in the direct dimension, in rad/s
- A: float Intensity
- states: bool Set to True for 'FnMODE': 'States-TPPI
- alt: bool Set to True for 'FnMODE': 'States-TPPI

#### Returns:

• S: 2darray
Gaussian function.

# $3.4.15 \quad sim.t\_2Dlorentzian(t1, t2, v1, v2, fwhm1, fwhm2, A=1, states=True, alt=True)$

Bidimensional lorentzian function.

#### Parameters:

- t1: *1darray*Indirect evolution timescale
- t2: *1darray*Timescale of the direct dimension
- v1: float
  Peak position in the indirect dimension, in Hz
- v2: float
  Peak position in the direct dimension, in Hz
- fwhm1: float Full-width at half maximum in the indirect dimension, in rad/s
- fwhm2: float
  Full-width at half maximum in the direct dimension, in rad/s
- A: float Intensity
- states: bool Set to True for 'FnMODE': 'States-TPPI
- alt: bool Set to True for 'FnMODE': 'States-TPPI

#### Returns:

• S: *2darray*Lorentzian function.

# $\begin{array}{ll} 3.4.16 & sim.t\_2Dpvoigt(t1,t2,v1,v2,fwhm1,fwhm2,A=1,b=0,states=True,\\ & alt=True) \end{array}$

Generates a 2D pseudo-voigt signal in the time domain. b states for the fraction of gaussianity, whereas A defines the overall amplitude of the total peak. Indexes '1' and '2' on the variables stand for 'F1' and 'F2', respectively.

#### Parameters:

- t1: 1darray
  Indirect evolution timescale
- t2: *1darray*Timescale of the direct dimension
- v1: float
  Peak position in the indirect dimension, in Hz
- v2: float
  Peak position in the direct dimension, in Hz
- $\bullet$  fwhm 1: float Full-width at half maximum in the indirect dimension, in rad/s
- $\bullet$  fwhm2: float Full-width at half maximum in the direct dimension, in rad/s
- A: float Intensity
- b: *float*Fraction of gaussianity
- states: bool Set to True for 'FnMODE': 'States-TPPI
- alt: bool Set to True for 'FnMODE': 'States-TPPI46

#### Returns:

• fid: 2darray
Pseudo-Voigt function.

# $3.4.17 \quad sim.t\_2Dvoigt(t1, t2, v1, v2, fwhm1, fwhm2, A=1, b=0, states=True, alt=True)$

Generates a 2D Voigt signal in the time domain. b states for the fraction of gaussianity, whereas A defines the overall amplitude of the total peak. Indexes '1' and '2' on the variables stand for 'F1' and 'F2', respectively.

#### Parameters:

- t1: *1darray*Indirect evolution timescale
- t2: *1darray*Timescale of the direct dimension
- v1: float
  Peak position in the indirect dimension, in Hz
- v2: float
  Peak position in the direct dimension, in Hz
- $\bullet$  fwhm 1: float Full-width at half maximum in the indirect dimension, in rad/s
- $\bullet$  fwhm2: float Full-width at half maximum in the direct dimension, in rad/s
- A: float Intensity
- b: *float*Fraction of gaussianity
- states: bool Set to True for 'FnMODE': 'States-TPPI
- alt: bool Set to True for 'FnMODE': 'States-TPPI

#### Returns:

• S: 2darray Voigt function.

# 3.4.18 sim.t\_gaussian(t, u, s, A=1, phi=0)

Gaussian function in the time domain.

#### Parameters:

- t: *1darray* Independent variable
- u: float Peak position, in Hz
- A: float Intensity
- phi: *float*Phase, in radians

#### Returns:

• S: *1darray* Gaussian function.

# $3.4.19 \quad sim.t\_lorentzian(t, u, fwhm, A=1, phi=0)$

Lorentzian function in the time domain.

#### Parameters:

- t: *1darray* Independent variable
- u: float Peak position, in Hz
- fwhm: float Full-width at half-maximum, in rad/s
- A: float Intensity
- phi: float
  Phase, in radians

#### Returns:

• S: *1darray*Lorentzian function.

# 3.4.20 sim.t\_pvoigt(t, u, fwhm, A=1, b=0, phi=0)

Pseudo-Voigt function in the time domain:

#### Parameters:

- t: *1darray* Independent variable
- u: float Peak position, in Hz
- fwhm: float Full-width at half-maximum, in rad/s
- A: float Intensity
- b: *float*Fraction of gaussianity
- phi: float
  Phase, in radians

#### Returns:

• S: 1darray Pseudo-Voigt function.

# $3.4.21 \quad sim.t\_voigt(t, u, fwhm, A=1, b=0, phi=0)$

Voigt function in the time domain. The parameter b affects the linewidth of the lorentzian and gaussian contributions.

#### Parameters:

- t: *1darray* Independent variable
- u: float Peak position, in Hz
- ullet fwhm: float Full-width at half-maximum, in rad/s
- A: float Intensity
- b: *float*Fraction of gaussianity
- phi: float
  Phase, in radians

#### Returns:

• S: *1darray* Voigt function.

## 3.4.22 sim.water7(N, t2, vW, fwhm=300, A=1, spread=701.125)

Simulates a feature like the water ridge in HSQC spectra, in the time domain.

#### Parameters:

- N: *int*Number of transients
- t2: *1darray*Time scale of the last temporal dimension.
- vW: float Nominal peak position, in Hz.
- fwhm: float Nominal full-width at half maximum of the peak, in rad/s.
- A: float
  Signal intensity.
- spread: *float* Standard deviation of the peak position distribution, in Hz.

#### Returns:

• ridge: *2darray*Matrix of the ridge.

class

# 3.5 FIT package

Functions for performing fits.

### 3.5.1 fit.CostFunc

Class that groups several ways to compute the target of the minimization in a fitting procedure. It includes the classic squared sum of the residuals, as well as some other non-quadratic cost functions. Let x be the residuals and s the chosen threshold value. Then the objective value R is computed as:

$$R = \sum \_i f(x\_i)$$

where f(x) can be chosen between the following options:

• Quadratic:

$$f(x) = x^2$$

• Truncated Quadratic:

$$f(x) = \begin{cases} x^2 & \text{if } |x| < s \\ s^2 & \text{otherwise} \end{cases}$$

• Huber function:

$$f(x) = \begin{cases} x^2 & \text{if } |x| < s \\ 2s|x| - s^2 & \text{otherwise} \end{cases}$$

• Asymmetric Truncated Quadratic:

$$f(x) = \begin{cases} x^2 & \text{if } x < s \\ s^2 & \text{otherwise} \end{cases}$$

• Asymmetric Huber function:

$$f(x) = \begin{cases} x^2 & \text{if } x < s \\ 2sx - s^2 & \text{otherwise} \end{cases}$$

#### Attributes:

- method: function

  Function to be used for the computation of the objective value. It must take as input the array of the residuals and the threshold, no matter if the latter is actually used or not.
- s: float
  Threshold value

#### Methods:

\_\_init\_\_(self, method='q', s=None)

Initialize the method according to your choice, then stores the threshold value in the attribute 's'. Allowed choices are:

- 'q': Quadratic
- 'tq': Truncated Quadratic
- 'huber': Huber function
- 'atq': Asymmetric Truncated Quadratic
- 'ahuber': Asymmetric Huber function

#### Parameters:

- method: strLabel for the method selection
- s: float
  Threshold value

## $\_$ call $\_$ (self, x)

Computes the objective value according to the chosen method and the residuals array x.

#### Parameters:

• x: 1darray
Array of the residuals

#### **Returns:**

• R: 1darray
Modified residuals according to the chosen target function

#### asymm huber(r, s)

Linear behaviour above s, penalizes negative entries

#### asymm truncated quadratic(r, s)

Constant behaviour above s, penalizes negative entries

#### huber(r, s)

Linear behaviour above s

#### method selector(self, method)

Performs the selection of the method according to the identifier string.

_						
П	_	 	 ~+	_	rc	

 $\bullet$  method: str Method label

#### Returns:

• f: function Selected model

## $squared\_sum(r,\,s{=}0)$

Quadratic everywhere

 $truncated\_quadratic(r, s)$ 

Constant behaviour above s

3.5.2 fit.Peak class

Class to represent the characteristic parameters of an NMR peak, and to compute it.

#### Attributes:

- t: 1darray
  Timescale for the FID
- SFO1: *float*Nucleus Larmor frequency
- o1p: float Carrier position
- N: *int*Number of points of the spectrum, i.e. after eventual zero-filling
- u: float Chemical shift /ppm
- fwhm: float Linewidth /Hz
- k: float
  Intensity, relative
- b: float
  Fraction of gaussianity (b=0 equals pure lorentzian)
- phi: float
  Phase /degrees
- group: *int* Identifier for the component of a multiplet

#### Methods:

 $\_\_init\_\_(self, acqus, u=None, fwhm=5, k=1, b=0, phi=0, N=None, group=0)$ 

Initialize the class with the configuration parameters, and with defauls values, if not given.

- acqus: dict
  It should contain 't', 'SFO1', 'o1p', and 'N'
- u: *float* Chemical shift /ppm
- fwhm: float Linewidth /Hz
- k: float
  Intensity, relative

- b: float
  Fraction of gaussianity (b=0 equals pure lorentzian)
- phi: float
  Phase /degrees
- N: int Number of points of the spectrum, i.e. after eventual zero-filling. None means to not zero-fill
- group: *int*Identifier for the component of a multiplet

Generates a voigt signal on the basis of the stored attributes, in the time domain. Then, makes the Fourier transform and returns it after the eventual zero-filling.

#### Parameters:

- A: *float*Absolute intensity value
- cplx: bool
  Returns the complex (True) or only the real part (False) of the signal
- get\_fid: bool

  If True, returns the FID instead of the transformed signal

#### **Returns:**

• sgn: *1darray* generated signal in the frequency domain

## get\_fid(self, A=1)

Compute and returns the FID encoding for that signal.

#### Parameters:

• A: float
Absolute intensity value

#### **Returns:**

• sgn: 1darray generated signal in the time domain

## par(self)

Creates a dictionary with the currently stored attributes and returns it.

#### **Returns:**

• dic: dict
Dictionary of parameters

## 3.5.3 fit.SINC ObjFunc

class

Computes the objective function as explained in M. Sawall et al., Journal of Magnetic Resonance 289 (2018), 132-141. The cost function is computed as:

$$f(d) = \sum_{i=1}^{3} \gamma_i g_i(d|e_i)$$

where d is the real part of the NMR spectrum.

#### Attributes:

- gamma1: float
  Weighting factor for function g<sub>1</sub>
- gamma2: float
  Weighting factor for function g<sub>2</sub>
- gamma3: floatWeighting factor for function  $g_3$
- e1: float
  Tolerance value for function g<sub>1</sub>
- e2: float
  Tolerance value for function g<sub>2</sub>

#### Methods:

 $\_\_init\_\_(self, gamma1=10, gamma2=0.01, gamma3=0, e1=0, e2=0)$ 

Initialize the coefficients used to weigh the objective function.

#### Parameters:

- gamma1: float
  Weighting factor for function g1
- gamma2: float
  Weighting factor for function g2
- gamma3: float
  Weighting factor for function g3
- e1: *float*Tolerance value for function g1
- e2: *float*Tolerance value for function g2

Computes the objective function f as explained in the paper

## g1(d, e1=0)

Penalty function for negative entries of the spectrum

## Parameters:

- d: *1darray* Spectrum
- e1: *float*Tolerance for negative entries

## g2(d, e2=0)

Regularization function that favours the smallest integral.

#### Parameters:

- d: *1darray* Spectrum
- e2: *float*Tolerance for ideal baseline

## **g3(d)**

Regularization function for the smoothing.

#### Parameters:

• d: *1darray* Spectrum

### 3.5.4 fit.Voigt Fit

class

This class offers an 'interface' to fit a 1D NMR spectrum.

#### Attributes:

- ppm\_scale: 1darray Self-explanatory
- S: *1darray* Spectrum to fit. Only real part
- t\_AQ: 1darray acquisition timescale of the spectrum
- SW: float Spectral width /Hz
- SFO1: *float*Larmor frequency of the nucleus
- o1p : float
  Pulse carrier frequency
- filename: strRoot of the names of the files that will be saved
- X\_label: str Label for the chemical shift axis in the figures
- i\_guess: *list*Initial guess for the fit, read by a .ivf file with fit.read vf
- result: *list*Result the fit, read by a .fvf file with fit.read vf

#### Methods:

\_\_init\_\_(self, ppm\_scale, S, t\_AQ, SFO1, o1p, nuc=None, filename='fit')

Initialize the class with common values.

- ppm\_scale: *1darray* ppm scale of the spectrum
- S: *1darray* Spectrum to be fitted
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: float
  Larmor frequency of the observed nucleus, in MHz

- o1p: float
  Carrier position, in ppm
- nuc: str Observed nucleus. Used to customize the x-scale of the figures.
- filename: str or None
  Root of the name of the files that will be saved

dofit(self, indep=True, u\_lim=1, f\_lim=10, k\_lim=(0, 3), vary\_phase=False, vary\_b=True, itermax=10000, fit\_tol=1e-08, filename=None, method='leastsq', basl\_fit='no')

Perform a lineshape deconvolution fitting. The initial guess is read from the attribute self.i\_guess. The components can be considered to be all independent from one to another by setting 'indep' to True: this means that the fit will be done using fit.voigt\_fit\_indep. The indep=False option has not been implemented yet.

- indep: bool

  True to consider all the components to be independent
- u\_lim: float

  Determines the displacement of the chemical shift (in ppm) from the starting value.
- f\_lim: float
  Determines the displacement of the linewidth (in Hz) from the starting value.
- k\_lim: float or tuple

  If tuple, minimum and maximum allowed values for k during the fit. If float, maximum displacement from the initial guess
- vary\_phase: bool Allow the peaks to change phase (True) or not (False)
- vary\_b: bool
  Allow the peaks to change Lorentzian/Gaussian ratio
- itermax: *int*Maximum number of allowed iterations
- fit\_tol: float
  Value of the target function to be set as x\_tol and f\_tol
- filename: strPath to the output file. If None, '<self.filename>.fvf' is used
- method: str or list of str Method to be used for the optimization. See lmfit for details. There is the option to run multiple optimizations in series.
- basl\_fit: str How to address the baseline fit. The options are:
  - 'no': Do not use baseline (default)

- 'fixed': The baseline is computed once and kept fixed during the optimization
- 'fit': The baseline coefficients enter as fit parameters during the nonlinear optimization
- 'calc': The baseline coefficients are calculated during the optimization via linear leastsquares optimization

#### **Returns:**

• lmfit\_results: list of lmfit.minimizer.MinimizerResult
Sequence of the fit results, ordered as the regions dictionary

#### get fit lines(self, what='result')

Calculates the components, and the total fit curve used as initial guess, or as fit results. The components will be returned as a list, not split by region.

#### Parameters:

• what: str 'iguess' or 'result'

#### **Returns:**

- signals: *list of 1darray*Components used for the fit
- total: *1darray*Sum of all the signals
- limits\_list: *list*List of region delimiters, in ppm
- whole\_basl: 1darray Computed baseline

#### iguess(self, filename=None, n=-1, ext='ivf', auto=False)

Reads, or computes, the initial guess for the fit. If the file is there already, it just reads it with fit.read\_vf. Otherwise, it calls fit.make\_iguess to make it.

- filename:  $str\ or\ None$ Path to the input file. If None, '<self.filename>.ivf' is used
- n: *int*Index of the initial guess to be read (default: last one)
- ext: strExtension of the file to be used
- auto: bool

  If True, uses the GUI for automatic peak picking, if False, the manual one

#### load fit(self, filename=None, n=-1, ext='fvf')

Reads a file with fit.read vf and stores the result in self.result.

#### Parameters:

- filename: strPath to the .fvf file to be read. If None, '<self.filename>.fvf' is used.
- n: *int*Index of the fit to be read (default: last one)
- ext: strExtension of the file to be used

plot(self, what='result', show\_total=True, show\_res=False, res\_offset=0, show\_basl=False, labels=None, filename=None, ext='tiff', dpi=600, dim=None)

Plots either the initial guess or the result of the fit, and saves all the figures. Calls fit.plot\_fit. The figure <filename>\_full will show the whole model and the whole spectrum. The figures labelled with \_R<k> will depict a detail of the fit in the k-th fitting region. Optional labels for the components can be given: in this case, the structure of 'labels' should match the structure of self.result (or self.i\_guess). This means that the length of the outer list must be equal to the number of fitting region, and the length of the inner lists must be equal to the number of peaks in that region.

- what: str 'iguess' to plot the initial guess, 'result' to plot the fitted data
- show\_total: bool Show the total trace (i.e. sum of all the components) or not
- show\_res: *bool*Show the plot of the residuals
- res\_offset: float
  Displacement of the residuals plot from 0, to be given as a fraction of the height of the experimental spectrum. res\_offset > 0 will move the residuals BELOW the zero-line!
- show\_basl: bool

  If True, displays the baseline on the spectrum and uses it to compute the total trace.
- labels: list of list
  Optional labels for the components. The structure of this parameter must match the structure
  of self.result
- filename: strRoot of the name of the figures that will be saved. If None,  $\langle self.filename \rangle$  is used
- ext: strFormat of the saved figures
- dpi: *int*Resolution of the figures, in dots per inches

• dim: tuple or None Dimensions of the figures to be saved, in inches. If None, the default value of (15, 8) is used.

 $\label{lem:condition} $$\operatorname{res\_histogram}(self, what='result', nbins=500, density=True, f\_lims=None, xlabel='Residuals x\_symm=True, barcolor='tab:green', fontsize=20, filename=None, ext='tiff', dpi=300)$ 

Computes the histogram of the residuals and saves it. Employs fit.histogram to make the figure.

#### Parameters:

• what: str 'iguess' or 'result'

• nbins : *int* number of bins to be calculated

• density : bool

True for normalize data

• f\_lims: tuple or None limits for the x axis of the figure

• xlabel: str or None

Text to be displayed under the x axis

• x\_symm : bool set it to True to make symmetric x-axis with respect to 0

• barcolor: strColor of the bins

• fontsize: *float*Biggest fontsize in the figure

• name: str name for the figure to be saved

• ext: strFormat of the image

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

## $to\_tragico(self, which='iguess', filename=None)$

Writes input 1 and input 2 for a TrAGICo run, on the basis of either the initial guess or the results of a fit. The files will be named '<filename>\_inp1' and '<filename>\_inp2', respectively.

#### Parameters:

• which: str
'iguess' or 'result'

• filename: strName of the file that will be saved. If None, the file will be saved in the spectrum directory

## 3.5.5 fit.Voigt Fit P2D

class

This class offers an 'interface' to fit a pseudo 2D NMR spectrum.

#### Attributes:

- ppm\_scale: 1darray Self-explanatory
- S: 2darray
  Spectrum to fit. Only real part
- t\_AQ: 1darray acquisition timescale of the spectrum
- SFO1: *float*Larmor frequency of the nucleus
- olp: float
  Pulse carrier frequency
- filename: strRoot of the names of the files that will be saved
- X\_label: str Label for the chemical shift axis in the figures
- i\_guess: *list*Initial guess for the fit, read by a .ivf file with fit.read vf P2D
- result: *list*Result the fit, read by a .fvf file with fit.read vf P2D

#### Methods:

 $\_\_init\_\_(self, ppm\_scale, S, t\_AQ, SFO1, o1p, nuc=None, filename='fit')$ 

Initialize the class with common values.

- ppm\_scale: *1darray* ppm scale of the spectrum
- S: 2darray
  Spectrum to be fitted
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: *float*Larmor frequency of the observed nucleus, in MHz
- o1p: float
  Carrier position, in ppm

• nuc: str

Observed nucleus. Used to customize the x-scale of the figures.

• filename: str or None

Root of the name of the files that will be saved

$$\label{local_control} \begin{split} & dofit(self,\ u\_tol=1,\ f\_tol=10,\ vary\_phase=False,\ vary\_b=True,\ itermax=10000,\ file-name=None) \end{split}$$

Perform a lineshape deconvolution fitting by calling fit.voigt\_fit\_P2D. The initial guess is read from the attribute self.i\_guess.

#### Parameters:

- u\_tol: *float*Determines the displacement of the chemical shift (in ppm) from the starting value.
- f\_tol: *float*Determines the displacement of the linewidth (in Hz) from the starting value.
- vary\_phase: bool
  Allow the peaks to change phase (True) or not (False)
- vary\_b: bool Allow the peaks to change Lorentzian/Gaussian ratio
- itermax: *int*Maximum number of allowed iterations
- filename: strPath to the output file. If None, '<self.filename>.fvf' is used

## get\_fit\_lines(self, what='result')

Calculates the components, and the total fit curve used as initial guess, or as fit results. The components will be returned as a list, not split by region.

#### Parameters:

• what: *str* 'iguess' or 'result'

#### Returns:

- signals: *list of list of 1darray* Components used for the fit
- total: *2darray* Sum of all the signals
- limits\_list: *list*List of the region delimiters, in ppm

#### iguess(self, input file=None, expno=0, n=-1)

Reads, or computes, the initial guess for the fit. If the file is there already, it just reads it with fit.read vf. Otherwise, it calls fit.make iguess to make it.

#### Parameters:

- input\_file: str or None
  Path to the input file. If None, '<self.filename>.ivf' is used
- expno: int

  Number of the experiment on which to compute the initial guess, in python numbering
- n: *int*Index of the initial guess to be read (default: last one)

#### load fit(self, output file=None, n=-1)

Reads a file with fit.read vf P2D and stores the result in self.result.

#### Parameters:

- output\_file: strPath to the .fvf file to be read. If None, '<self.filename>.fvf' is used.
- n: *int*Index of the fit to be read (default: last one)

 $plot(self, what='result', show\_total=True, show\_res=False, res\_offset=0, labels=None, filename=None, ext='tiff', dpi=600)$ 

Plots either the initial guess or the result of the fit, and saves all the figures. Calls fit.plot\_fit\_P2D. The figures <filename>\_full will show the whole model and the whole spectrum. The figures labelled with \_R<k> will depict a detail of the fit in the k-th fitting region. Optional labels for the components can be given: in this case, the structure of 'labels' should match the structure of self.result (or self.i\_guess). This means that the length of the outer list must be equal to the number of fitting region, and the length of the inner lists must be equal to the number of peaks in that region.

- what: str 'iguess' to plot the initial guess, 'result' to plot the fitted data
- show\_total: bool
  Show the total trace (i.e. sum of all the components) or not
- show\_res: bool
  Show the plot of the residuals
- res\_offset: float
  Displacement of the residuals plot from 0, to be given as a fraction of the height of the experimental spectrum. res\_offset > 0 will move the residuals BELOW the zero-line!

- labels: list of list
  Optional labels for the components. The structure of this parameter must match the structure
  of self.result
- $\bullet$  filename: strRoot of the name of the figures that will be saved. If None, <self.filename> is used
- ullet ext: str Format of the saved figures
- dpi: *int*Resolution of the figures, in dots per inches

res\_histogram(self, what='result', nbins=500, density=True, f\_lims=None, xlabel='Residuals x symm=True, barcolor='tab:green', fontsize=20, filename=None, ext='tiff', dpi=300)

Computes the histogram of the residuals and saves it in the same folder of the fit figures. Employs fit.histogram to make the figure.

- what: str 'iguess' or 'result'
- nbins : *int* number of bins to be calculated
- density : *bool*True for normalize data
- f\_lims: tuple or None limits for the x axis of the figure
- xlabel : str or None
  Text to be displayed under the x axis
- x\_symm : bool set it to True to make symmetric x-axis with respect to 0
- barcolor: strColor of the bins
- fontsize: *float*Biggest fontsize in the figure
- name : str name for the figure to be saved
- ext: strFormat of the image
- dpi: *int*Resolution of the image in dots per inches

3.5.6 fit.ax\_histogram(ax, data0, nbins=100, density=True, f\_lims=None, xlabel=None, x\_symm=False, fitG=True, barcolor='tab:blue', font-size=10)

Computes an histogram of 'data' and tries to fit it with a gaussian lineshape. The parameters of the gaussian function are calculated analytically directly from 'data' using 'scipy.stats.norm'

#### Parameters:

• ax : matplotlib.subplot Object panel of the figure where to put the histogram

• data0 : ndarray the data to be binned

• nbins: *int* number of bins to be calculated

• density : *bool*True for normalize data

• f\_lims: tuple or None limits for the x axis of the figure

• xlabel: str or None

Text to be displayed under the x axis

• x\_symm : bool set it to True to make symmetric x-axis with respect to 0

• fitG: bool
Shows the gaussian approximation

• barcolor: strColor of the bins

• fontsize: *float*Biggest fontsize in the figure

#### Returns:

 $\bullet$  m : float Mean of data

• s : float Standard deviation of data.

## 3.5.7 fit.bin data(data0, nbins=100, density=True, x symm=False)

Computes the histogram of data, sampling it into nbins bins.

#### Parameters:

• data : *ndarray* the data to be binned

• nbins : *int* number of bins to be calculated

 $\bullet$  density : boolTrue for normalize data

• x\_symm : bool set it to True to make symmetric x-axis with respect to 0

#### Returns:

• hist: *1darray*The bin intensity

• bin\_scale: *1darray*Scale built with the mean value of the bin widths.

## 3.5.8 fit.build 2D sgn(parameters, acqus, N=None, procs=None)

Create a 2D signal according to the final parameters returned by make\_iguess\_2D. Process it according to procs.

#### Parameters:

- parameters: *list or 2darray* sequence of the parameters: u1, u2, fwhm1, fwhm2, I, b. Multiple components are allowed
- acqus: dict 2D-like acqus dictionary containing the acquisition timescales (keys t1 and t2)
- N: tuple of int Zero-filling values (F1, F2). Read only if procs is None
- procs: *dict* 2D-like procs dictionary.

#### Returns:

• peak: 2darray rr part of the generated signal

## 3.5.9 fit.build baseline(ppm scale, C, L=None)

Builds the baseline calculating the polynomion with the given coefficients, and summing up to the right position.

#### Parameters:

- ppm\_scale: *1darray* ppm scale of the spectrum
- C: *list*Parameters coefficients. No baseline corresponds to False.
- L: *list*List of window regions. If it is None, the baseline is built on the whole ppm\_scale

#### Returns:

• baseline: *1darray* Self-explanatory.

# $3.5.10 \quad fit.calc\_R2(y,\,y\_c)$

Computes the R-squared coefficient of a linear regression as:

$$R^{2} = 1 - \frac{\sum (y - y_{mean})^{2}}{\sum (y - y_{c})^{2}}$$

## Parameters:

- y: *1darray* Experimental data
- y\_c: *1darray* Calculated data

#### Returns:

• R2: float R-squared coefficient

## 3.5.11 fit.calc fit lines(ppm scale, limits, t AQ, SFO1, o1p, N, V, C=False)

Given the values extracted from a fit input/output file, calculates the signals, the total fit function, and the baseline.

#### Parameters:

- ppm\_scale: 1darray PPM scale of the spectrum
- limits: *tuple* (left, right) in ppm
- t\_AQ: 1darray Acquisition timescale
- SFO1: *float*Larmor frequency of the nucleus /ppm
- o1p: float
  Pulse carrier frequency /ppm
- N: *int* Size of the final spectrum.
- V: 2darray
  Matrix containing the values to build the signals.
- C: *1darray*Baseline polynomion coefficients. False to not use the baseline

#### Returns:

- sgn: *list* Voigt signals built using V
- Total: *1darray* sum of all the sgn
- $\bullet$  baseline: 1darray

Polynomion built using C. False if C is False.

# $3.5.12 \quad fit.dic2mat(dic, peak\_names, ns, A=None)$

This is used to make the matrix of the parameters starting from a dictionary like the one produced by l. The column of the total intensity is not added, unless the parameter 'A' is passed. In this case, the third column (which is the one with the relative intesities) is corrected using the function molfrac.

## Parameters:

• dic : dict input dictionary

• peak\_names : *list* list of the parameter entries to be looked for

• ns: int number of signals to unpack

• A : float or None Total intensity.

## Returns:

• V : 2darray
Matrix containing the parameters.

# $3.5.13 \quad fit.fit\_int(y,\,y\_c,\,q{=}True)$

Computes the optimal intensity and intercept of a linear model in the least squares sense. Let y be the experimental data and y\_c the model, and let <w> the mean of variable w. Then: A = ( <y\_c y> - <y\_c><y> ) / ( <y\_c2^> - <y\_c>^2 ) q = ( <y\_c>^2 <y> - <y\_c><y\_c y> ) / ( <y\_c2^> + <y\_c^2 > - <y\_c>^2 )

## Parameters:

- y: 1darray Experimental data
- y\_c: 1darray Model data
- q: bool

  If True, includes the offset in the calculation. If False, only the intensity factor is computed.

- A: float
  Optimized intensity
- q: float Optimized intercept

# 3.5.14 fit.gaussian\_fit(x, y, s\_in=None)

Fit 'y' with a gaussian function, built using 'x' as independent variable

# Parameters:

- x : 1darray x-scale
- y : 1darray data to be fitted

- ullet u : float mean
- $\bullet$  s : float standard deviation
- ullet A: float Integral

# $3.5.15 \quad \text{fit.gen\_iguess} (x, \text{experimental}, \text{param}, \text{model}, \text{model\_args} = [], \text{sens0} = 1)$

GUI for the interactive setup of a Parameters object to be used in a fitting procedure. Once you initialized the Parameters object with the name of the parameters and a dummy value, you are allowed to set the value, minimum, maximum and vary status through the textboxes given in the right column, and see their effects in real time. Upon closure of the figure, the Parameters object with the updated entries is returned.

Keybinding:

- '>': increase sensitivity
- '<': decrease sensitivity
- 'up': increase value
- 'down': decrease value
- 'left': change parameter
- 'right': change parameter
- 'v': change 'vary' status
- '<': toggle automatic zoom adjustment

#### **Parameters:**

- x: 1darray Independent variable
- experimental: *1darray*The objective values you are trying to fit
- param: *lmfit.Parameters Object* Initialized parameters object
- model: function Function to be used for the generation of the fit model. Param must be the first argument.
- model\_args: *list*List of args to be passed to model, after param
- sens0: *float*Default sensitivity for the change of the parameters with the mouse

## Returns:

• param: *lmfit.Parameters Object* Updated Parameters Object

# 3.5.16 fit.gen\_iguess\_2D(ppm\_f1, ppm\_f2, tr1, tr2, u1, u2, acqus, fwhm0=100 procs=None)

Generate the initial guess for the fit of a 2D signal. The employes model is the one of a 2D Voigt signal, acquired with the States-TPPI scheme in the indirect dimension (i.e. sim.t\_2DVoigt). The program allows for the inclusion of up to 10 components for the signal, in order to improve the fit. The acqus dictionary must contain the following keys: > t1: acquisition timescale in the indirect dimension (States) > t2: acquisition timescale in the direct dimension > SFO1: Larmor frequency of the nucleus in the indirect dimension > o1p: carrier position in the indirect dimension /ppm > o2p: carrier position in the direct dimension /ppm The signals will be processed according to the values in the procs dictionary, if given; otherwise, they will be just zero-filled up to the data size (i.e. (len(ppm\_f1), len(ppm\_f2))).

## Parameters:

- ppm\_f1: *1darray* ppm scale for the indirect dimension
- ppm\_f2: 1darray ppm scale for the direct dimension
- tr1: 1darray
  Trace of the original 2D peak in the indirect dimension
- tr2: *1darray*Trace of the original 2D peak in the direct dimension
- u1: float Chemical shift of the original 2D peak in the indirect dimension /ppm
- u2: float
  Chemical shift of the original 2D peak in the direct dimension /ppm
- acqus: *dict*Dictionary of acquisition parameters
- fwhm0: float
  Initial value for FWHM in both dimensions
- procs: *dict*Dictionary of processing parameters

- final\_parameters: 2darray
  Matrix of dimension (# signals, 6) that contains, for each row: v1(Hz), v2(Hz), fwhm1(Hz),
  fwhm2(Hz), A, b
- fit\_interval: tuple of tuple Fitting window. ( (left\_f1, right\_f1), (left\_f2, right\_f2) )

# 3.5.17 fit.get region(ppmscale, S, rev=True)

Interactively select the spectral region to be fitted. Returns the border ppm values.

## Parameters:

- ppmscale: *1darray*The ppm scale of the spectrum
- S: *1darray*The spectrum to be trimmed
- rev: bool Choose if to reverse the ppm scale and data (True) or not (False).

- left: *float*Left border of the selected spectral window
- right: *float*Right border of the selected spectral window

3.5.18 fit.histogram(data, nbins=100, density=True, f\_lims=None, xlabel=None, x\_symm=False, fitG=True, barcolor='tab:blue', font-size=10, name=None, ext='tiff', dpi=600)

Computes an histogram of 'data' and tries to fit it with a gaussian lineshape. The parameters of the gaussian function are calculated analytically directly from 'data' using 'scipy.stats.norm'

#### Parameters:

• data : ndarray the data to be binned

• nbins : *int* number of bins to be calculated

• density : bool

True for normalize data

• f\_lims: tuple or None limits for the x axis of the figure

• xlabel : str or None
Text to be displayed under the x axis

• x\_symm : bool set it to True to make symmetric x-axis with respect to 0

• fitG: bool
Shows the gaussian approximation

• barcolor: str Color of the bins

• fontsize: *float*Biggest fontsize in the figure

• name : str name for the figure to be saved

• ext: strFormat of the image

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

#### **Returns:**

 $\bullet$  m : float Mean of data

• s : float
Standard deviation of data.

# 3.5.19 fit.integrate(ppm0, data0, X label=)

Allows interactive integration of a NMR spectrum through a dedicated GUI. Returns the values as a dictionary, where the keys are the selected regions truncated to the 2nd decimal figure. The returned dictionary contains pre-defined keys, as follows:

- total: total integrated area
- ref\_pos: location of the reference peak /ppm1:ppm2
- ref int: absolute integral of the reference peak
- ref\_val: for how many nuclei the reference peak integrates

The absolute integral of the x-th peak, I x, must be calculated according to the formula:

#### Parameters:

• ppm: *1darray* PPM scale of the spectrum

• data: *1darray* Spectrum to be integrated.

• X\_label: str Label of the x-axis

#### **Returns:**

• f\_vals: dict Dictionary containing the values of the integrated peaks.

# 

Function to select and integrate 2D peaks of a spectrum, using dedicated GUIs. Calls integral\_2D to do the dirty job.

## **Parameters:**

- ppm\_f1: 1darray PPM scale of the indirect dimension
- ppm\_f2: *1darray* PPM scale of the direct dimension
- data: *2darray* real part of the spectrum
- SFO1: float
  Larmor frequency of the nucleus in the indirect dimension
- SFO2: float
  Larmor frequency of the nucleus in the direct dimension
- fwhm\_1: float Starting FWHM /Hz in the indirect dimension
- fwhm\_2: float Starting FWHM /Hz in the direct dimension
- utol\_1: float
  Allowed tolerance for u 1 during the fit. (u 1-utol 1, u 1+utol 1)
- utol\_2: float Allowed tolerance for u\_2 during the fit. (u\_2-utol\_2, u\_2+utol\_2)
- plot\_result: bool

  True to show how the program fitted the traces.

#### **Returns:**

• I: dict Computed integrals. The keys are '<ppm f1>:<ppm f2>' with 2 decimal figures.

# 3.5.21 fit.interactive smoothing(x, y, cmap='RdBu')

Interpolate the given data with a 3rd-degree spline. Type the desired smoothing factor in the box and see the outcome directly on the figure. When the panel is closed, the smoothed function is returned.

# Parameters:

- x: 1darray
  Scale of the data
- y: *1darray*Data to be smoothed
- $\bullet$  cmap: strName of the colormap to be used to represent the weights

- sx: *1darray*Location of the spline points
- sy: 1darray Smoothed y
- s\_f: *float*Employed smoothing factor for the spline
- weights: *1darray* Weights vector

# 3.5.22 fit.join par(filenames, ppm scale, joined name=None)

Load a series of parameters fit files. Join them together, returning a unique array of signal parameters, a list of coefficients for the baseline, and a list of tuples for the regions. Also, uses the coefficients and the regions to directly build the baseline according to the ppm windows.

## Parameters:

- filenames: *list*List of directories of the input files.
- ppm\_scale: *1darray* ppm scale of the spectrum. Used to build the baseline
- joined\_name: str or None If it is not None, concatenates the files in the list 'filenames' and saves them in a single file named 'joined\_name'.

- V: 2darray
  Array of joined signal parameters
- C: *list*Parameters coefficients. No baseline corresponds to False.
- L: *list*List of window regions.
- baseline: *1darray*Baseline built from C and L.

# 3.5.23 fit.lr(y, x=None, force intercept=False)

Performs a linear regression of y with a model  $y_c = mx + q$ .

## Parameters:

- y: *1darray*Data to be fitted
- x: 1darray
  Independent variable. If None, the point indexes are used.
- force\_intercept: bool
  If True, forces the intercept to be zero.

- y\_c: 1darray Fitted trend
- values: tuple (m, q)

# 3.5.24 fit.lsp(y, x, n=5, w=None)

Linear-System Polynomion Make a polynomial fit on the experimental data y by solving the linear system

$$\mathbf{y}=\mathbb{T}\,\mathbf{c}$$

where  $\mathbb{T}$  is the Vandermonde matrix of the x-scale and  $\mathfrak{c}$  is the set of coefficients that minimize the problem in the least-squares sense. It is also possible to make it weighted by using an array of weights w.

## Parameters:

- y: 1darray Experimental data
- x: 1darray
  Independent variable (better if normalized)
- n: intOrder of the polynomion + 1, i.e. number of coefficients
- w: 1darray
  Array of weights for the data. If None, the nonweighted approach is used

#### Returns:

• c: 1darray
Set of minimized coefficients

# 3.5.25 fit.make\_iguess(S\_in, ppm\_scale, t\_AQ, SFO1=701.125, o1p=0, filename='i\_guess')

Creates the initial guess for a lineshape deconvolution fitting procedure, using a dedicated GUI. The GUI displays the experimental spectrum in black and the total function in blue. First, select the region of the spectrum you want to fit by focusing the zoom on it using the lens button. Then, use the '+' button to add components to the spectrum. The black column of text under the textbox will be colored with the same color of the active peak. Use the mouse scroll to adjust the parameters of the active peak. Write a number in the 'Group' textbox to mark the components of the same multiplet. Group 0 identifies independent peaks, not part of a multiplet (default). The sensitivity of the mouse scroll can be regulated using the 'up arrow' and 'down arrow' buttons. The active peak can be changed in any moment using the slider.

The baseline can be computed first by initializing the x-scale on the selected window through the 'SET BASL' button. The informer light next to the button becomes green if it is properly set. The baseline coefficients can be set with the mouse scroll analogously to any other parameter.

When you are satisfied with your fit, press 'SAVE' to write the information in the output file. Then, the GUI is brought back to the initial situation, and the region you were working on will be marked with a green rectangle. You can repeat the procedure as many times as you wish, to prepare the guess on multiple spectral windows.

Keyboard shortcuts:

- 'increase sensitivity' : '>'
- 'decrease sensitivity' : '<'
- mouse scroll up: 'up arrow key'
- mouse scroll down: 'down arrow key'
- 'add a component': '+'
- 'remove the active component': '-'
- 'change component, forward': 'page up'
- 'change component, backward': 'page down'

#### **Parameters:**

- S\_in: *1darray* Experimental spectrum
- ppm\_scale: 1darray
  PPM scale of the spectrum
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: float Nucleus Larmor frequency /MHz
- o1p: float
  Carrier frequency /ppm
- filename: strPath to the filename where to save the information. The '.ivf' extension is added automatically.

# 3.5.26 fit.make\_iguess\_P2D(S\_in, ppm\_scale, expno, t\_AQ, SFO1=701.125, o1p=0, filename='i guess')

Creates the initial guess for a lineshape deconvolution fitting procedure of a pseudo-2D experiment, using a dedicated GUI. It will be donw on only one experiment of the whole pseudo-2D. The GUI displays the experimental spectrum in black and the total function in blue. First, select the region of the spectrum you want to fit by focusing the zoom on it using the lens button. Then, use the '+' button to add components to the spectrum. The black column of text under the textbox will be colored with the same color of the active peak. Use the mouse scroll to adjust the parameters of the active peak. Write a number in the 'Group' textbox to mark the components of the same multiplet. Group 0 identifies independent peaks, not part of a multiplet (default). The sensitivity of the mouse scroll can be regulated using the 'up arrow' and 'down arrow' buttons. The active peak can be changed in any moment using the slider.

When you are satisfied with your fit, press 'SAVE' to write the information in the output file. Then, the GUI is brought back to the initial situation, and the region you were working on will be marked with a green rectangle. You can repeat the procedure as many times as you wish, to prepare the guess on multiple spectral windows.

Keyboard shortcuts:

- 'increase sensitivity' : '>'
- 'decrease sensitivity' : '<'
- mouse scroll up: 'up arrow key'
- mouse scroll down: 'down arrow key'
- 'add a component': '+'
- 'remove the active component': '-'
- 'change component, forward': 'page up'
- 'change component, backward': 'page down'

#### **Parameters:**

- S\_in: 1darray Experimental spectrum
- ppm\_scale: *1darray* PPM scale of the spectrum
- $\bullet$  expno: int Index of experiment of the pseudo 2D on which to compute the initial guess, in python numbering
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: *float* Nucleus Larmor frequency /MHz
- o1p: float
  Carrier frequency /ppm
- filename: strPath to the filename where to save the information. The '.ivf' extension is added automatically.

# 3.5.27 fit.make iguess auto(ppm, data, SW, SFO1, o1p, filename='iguess')

GUI to create a .ivf file, used as initial guess for Voigt\_Fit. The computation of the peak positions and linewidths employs scipy.signal.find\_peaks and scipy.signal.peak\_widths, respectively. In addition, peak features may be added manually by clicking with the left button twice. Unwanted features can be removed with right clicks. If the FWHM of a peak cannot be computed automatically, a dummy FWHM of 1 Hz is assigned automatically. The file <filename>.ivf is written upon pressing the SAVE button. Press Z to activate/deactivate the cursor snap.

## Parameters:

• ppm: *1darray* PPM scale of the spectrum

• data: *1darray* real part of the spectrum to fit

• SW: float Spectral width /Hz

• SFO1: *float* Nucleus Larmor Frequency /MHz

• o1p: float
Carrier position /ppm

• filename: strPath to the file where to save the initial guess. The .ivf extension is added automatically.

# 3.5.28 fit.make signal(t, u, s, k, b, phi, A, SFO1=701.125, o1p=0, N=None)

Generates a voigt signal on the basis of the passed parameters in the time domain. Then, makes the Fourier transform and returns it.

#### Parameters:

- t : ndarray acquisition timescale
- u : *float* chemical shift /ppm
- s : *float* full-width at half-maximum /Hz
- k : float relative intensity
- b : *float* fraction of gaussianity
- phi : *float* phase of the signal, in degrees
- A: float total intensity
- SFO1 : float Larmor frequency /MHz
- o1p : float pulse carrier frequency /ppm
- N: int or None length of the final signal. If None, signal is not zero-filled before to be transformed.

#### Returns:

• sgn: *1darray* generated signal in the frequency domain

# 3.5.29 fit.peak pick(ppm f1, ppm f2, data, coord filename='coord.tmp')

Make interactive peak\_picking. The position of the selected signals are saved in coord\_filename. If coord\_filename already exists, the new signals are appended at its bottom: nothing is overwritten. Calls misc.select\_traces for the selection.

# Parameters:

- ppm\_f1: *1darray* ppm scale for the indirect dimension
- ppm\_f2: *1darray* ppm scale for the direct dimension
- data: 2darray
  Spectrum to peak-pick. The dimension should match the scale sizes.
- coord\_filename: str
  Path to the file where to save the peak coordinates

#### Returns:

• coord: *list*List of (u2, u1) for each peak

3.5.30 fit.plot\_fit(S, ppm\_scale, regions, t\_AQ, SFO1, o1p, show\_total=False, show\_res=False, res\_offset=0, show\_basl=False, X\_label='\$ delta\$ /ppm', labels=None, filename='fit', ext='tiff', dpi=600, dim=None)

Plots either the initial guess or the result of the fit, and saves all the figures. The figure <file-name>\_full will show the whole model and the whole spectrum. The figures labelled with  $_{\rm R}<$ k> will depict a detail of the fit in the k-th fitting region. Optional labels for the components can be given: in this case, the structure of 'labels' should match the structure of 'regions'. This means that the length of the outer list must be equal to the number of fitting region, and the length of the inner lists must be equal to the number of peaks in that region.

#### **Parameters:**

- S: *1darray*Spectrum to be fitted
- ppm\_scale: 1darray ppm scale of the spectrum
- regions: dict Generated by fit.read vf
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: *float*Larmor frequency of the observed nucleus, in MHz
- o1p: float
  Carrier position, in ppm
- show\_total: bool Show the total trace (i.e. sum of all the components) or not
- show\_res: bool
  Show the plot of the residuals
- res\_offset: float
  Displacement of the residuals plot from 0, to be given as a fraction of the height of the experimental spectrum. res\_offset > 0 will move the residuals BELOW the zero-line!
- show\_basl: bool

  If True, displays the baseline on the spectrum and uses it to compute the total trace.
- X\_label: str Text to show as label for the chemical shift axis
- labels: *list of list*Optional labels for the components. The structure of this parameter must match the structure of self.result
- ullet filename: str Root of the name of the figures that will be saved. If None, <self.filename> is used

ullet ext: str Format of the saved figures

ullet dpi: int Resolution of the figures, in dots per inches

  $3.5.31 \quad fit.plot\_fit\_P2D(S,ppm\_scale,regions,t\_AQ,SFO1,o1p,show\_total=\\ show\_res=False,res\_offset=0,X\_label=,labels=None,filename='fit',\\ ext='tiff',dpi=600)$ 

Plots either the initial guess or the result of the fit, and saves all the figures. A new folder named <filename>\_fit will be created. The figure <filename>\_full will show the whole model and the whole spectrum. The figures labelled with  $_R<$ k> will depict a detail of the fit in the k-th fitting region. Optional labels for the components can be given: in this case, the structure of 'labels' should match the structure of 'regions'. This means that the length of the outer list must be equal to the number of fitting region, and the length of the inner lists must be equal to the number of peaks in that region.

## Parameters:

- S: 2darray
  Spectrum to be fitted
- ppm\_scale: *1darray* ppm scale of the spectrum
- regions: *list of dict* Generated by fit.read\_vf\_P2D
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: float
  Larmor frequency of the observed nucleus, in MHz
- o1p: float
  Carrier position, in ppm
- nuc: str Observed nucleus. Used to customize the x-scale of the figures.
- show\_total: bool Show the total trace (i.e. sum of all the components) or not
- show\_res: *bool*Show the plot of the residuals
- res\_offset: float
  Displacement of the residuals plot from 0, to be given as a fraction of the height of the experimental spectrum. res\_offset > 0 will move the residuals BELOW the zero-line!
- X\_label: str Text to show as label for the chemical shift axis
- labels: list of list
  Optional labels for the components. The structure of this parameter must match the structure
  of self.result
- filename: strRoot of the name of the figures that will be saved.

ullet dpi: int Resolution of the figures, in dots per inches

# 3.5.32 fit.polyn\_basl(y, n=5, method='huber', s=0.2, c\_i=None, itermax=1000)

Fit the baseline of a spectrum with a low-order polynomion using a non-quadratic objective function. Let y be an array of N points. The polynomion is generated on a normalized scale that goes from -1 to 1 in N steps, and the coefficients are initialized either from outside through the parameter c\_i or with the ordinary least squares fit. Then, the guess is refined using the objective function of choice employing the trust-region reflective least-squares algorithm.

#### Parameters:

- y: 1darray Experimental data
- n: intOrder of the polynomion + 1, i.e. number of coefficients
- method: str Objective function of choice. 'q': quadratic, 'tq': truncated quadratic, 'huber': Huber, 'atq': asymmetric truncated quadratic, 'ahuber': asymmetric huber
- s: float
  Relative threshold value for the non-quadratic behaviour of the objective function
- c\_i: sequence or None Initial guess for the polynomion coefficient. If None, the least-squares fit is used
- itermax: *int*Number of maximum iterations

- px: *1darray*Fitted polynomion
- c: *list*Set of coefficients of the polynomion

# 3.5.33 fit.print\_par(V, C, limits=[None, None])

Prints on screen the same thing that write\_par writes in a file.

## Parameters:

- V : 2darray matrix (# signals, parameters)
- C: 1darray or False
  Coefficients of the polynomion to be used as baseline correction. If the 'baseline' checkbox in the interactive figure panel is not checked, C\_f is False.
- limits: tuple or None
  Trim limits for the spectrum (left, right). If None, the whole spectrum is used.

# 3.5.34 fit.read par(filename)

Reads the input file of the fit and returns the values.

## Parameters:

• filename: str directory and name of the input file to be read

- V : 2darray matrix (# signals, parameters)
- C: 1darray or False

  Coefficients of the polynomion to be used as baseline correction. If the 'baseline' checkbox in the interactive figure panel is not checked, C\_f is False.
- limits: tuple or None
  Trim limits for the spectrum (left, right). If None, the whole spectrum is used.

# 3.5.35 fit.read\_vf(filename, n=-1)

Reads a .ivf (initial guess) or .fvf (final fit) file, containing the parameters for a lineshape deconvolution fitting procedure. The file is separated and unpacked into a list of dictionaries, each of which contains the limits of the fitting window, the total intensity value, and a dictionary for each peak with the characteristic values to compute it with a Voigt line.

#### **Parameters:**

- filename: strPath to the filename to be read
- n: *int*Number of performed fit to be read. Default: last one. The breakpoints are lines that start with '!'. For this reason, n=0 returns an empty dictionary, hence the first fit is n=1.

# **Returns:**

• regions: *list*List of dictionaries for running the fit.

# 3.5.36 fit.read vf P2D(filename, n=-1)

Reads a .ivf (initial guess) or .fvf (final fit) file, containing the parameters for a lineshape deconvolution fitting procedure. The file is separated and unpacked into a list of list of dictionaries, each of which contains the limits of the fitting window, and a dictionary for each peak with the characteristic values to compute it with a Voigt line.

## Parameters:

- filename: strPath to the filename to be read
- n: int
  Number of performed fit to be read. Default: last one. The breakpoints are lines that start with '!'. For this reason, n=0 returns an empty dictionary, hence the first fit is n=1.

# **Returns:**

• regions: *list of list of dict*List of dictionaries for running the fit.

# 3.5.37 fit.sinc\_phase(data, gamma1=10, gamma2=0.01, gamma3=0, e1=0, e2=0, \*\*fit kws)

Perform automatic phase correction according to the SINC algorithm, as described in M. Sawall et. al., Journal of Magnetic Resonance 289 (2018), 132–141. The fitting method defaults to 'least\_squares'.

## Parameters:

- data: *1darray* Spectrum to phase-correct
- gamma1: float
  Weighting factor for function g1: non-negativity constraint
- gamma2: float
  Weighting factor for function g2: smallest-integral constraint
- gamma3: float
  Weighting factor for function g3: smoothing constraint
- e1: *float*Tolerance factor for function g1: adjustment for noise
- e2: float
  Tolerance factor for function g2: adjustment for non-ideal baseline
- fit\_kws: keyworded arguments additional parameters for the fit function. See lmfit.Minimizer.minimize for details. Do not use 'leastsq' because the cost function returns a scalar value!

- p0: float
  Fitted zero-order phase correction angle, in degrees
- p1: float
  Fitted first-order phase correction angle, in degrees

# 3.5.38 fit.smooth\_spl(x, y, s\_f=1, size=0, weights=None)

Fit the input data with a 3rd-order spline, given the smoothing factor to be applied.

## Parameters:

- x: 1darray
  Location of the experimental points
- y: 1darray Input data to be fitted
- s\_f: float Smoothing factor of the spline. 0=best straight line, 1=native spline.
- size: *int*Size of the spline. If size=0, the same dimension as y is chosen.

- x\_s: 1darray Location of the spline data points.
- y\_s: 1darray Spline that fits the data.

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# 3.5.39 fit.test correl(data, subtract mean=True)

Tests an array of residuals for their correlation. It compares the unit-lag autocorrelation P of the data (see below) with the theoretical value for non-correlated data  $T_P$ :

$$P = \sum_{i} \mathbb{r}[i+1]; T_P = (N-1)^{(0.5)} \sum_{i} \mathbb{r}[i]^2$$

If  $P < T_P$ , the residuals are not correlated, and the result is True.

## Parameters:

- data: *1darray*Residuals to be test
- subtract\_mean: bool
  If True, subtracts from the residuals their mean.

# **Returns:**

• test: bool

True if the residuals are non correlated, False otherwise

# 3.5.40 fit.test ks(data, thresh=0.05)

Performs the Kolmogorov-Smirnov test on the residuals to check if they are drawn from a normal distribution. The implementation is scipy.stats.kstest. The result is True if the residuals are Gaussian.

## Parameters:

• data: *1darray* Residuals to test

 $\bullet$  thresh: float

Significance level for the test. Default is 5

# Returns:

• test: bool

True if the residuals are Gaussian, False otherwise

# 3.5.41 fit.test randomsign(data, thresh=1.96)

Test an array of residuals for the randomness of the sign changes. The result it True if the sequence is recognized as random.

## Parameters:

• data: *1darray* Residuals to test

• thresh: *float* Significance level. The default is 1.96, which corresponds to 5

# Returns:

• test: bool

True if the signs are random, False otherwise

# 3.5.42 fit.test residuals(res, alpha=0.05)

Tests an array of residuals for their randomness, correlation, and underlying distribution. To do this, it uses the functions 'fit.test\_randomsign', 'fit.test\_correl', 'fit.test\_ks'. The results of the tests will be print in standard output and returned.

## Parameters:

• res: *ndarray*Residuals to be tested

• alpha: *float* Significance level

## Returns:

• test\_random: bool
Randomness of the residuals (True = random)

• test\_correlation: bool Correlation of the residuals (True = non-correlated)

• test\_gaussian: bool
Normal-distribution of the residuals (True = normally-distributed)

 $\begin{array}{lll} 3.5.43 & \text{fit.voigt\_fit}(S, ppm\_scale, V, C, t\_AQ, limits=None, SFO1=701.125, \\ & o1p=0, utol=0.5, vary\_phi=False, vary\_xg=True, hist\_name=None, \\ & \text{write out='fit.out', test res}=True) \end{array}$ 

Fits an NMR spectrum with a set of signals, whose parameters are specifed in the V matrix. There is the possibility to use a baseline through the parameter C. The signals are computed in the time domain and then Fourier transformed.

#### **Parameters:**

- S: 1darray
  Spectrum to be fitted
- ppm\_scale : 1darray Self-explanatory
- V : 2darray matrix (# signals, parameters)
- C: 1darray or False
  Coefficients of the polynomion to be used as baseline correction. If it is False, the baseline correction is not used.
- t\_AQ : 1darray Acquisition timescale
- limits: tuple or None
  Trim limits for the spectrum (left, right). If None, the whole spectrum is used.
- SFO1: float Larmor frequency /MHz
- o1p: float pulse carrier frequency /ppm
- utol: float tolerance for the chemical shift. The peak center can move in the range  $[\mu utol, \mu + utol]$ .
- vary\_xg: bool

  If it is False, the parameter x\_g cannot be varied during the fitting procedure. Useful when fitting with pure Gaussians or pure Lorentzians.
- vary\_basl: bool

  If it is False, the baseline is kept fixed at the initial parameters.

- C\_f: 1darray or False
  Coefficients of the polynomion to be used as baseline correction, or just False if not used.
- V\_f: 2darray matrix (# signals, parameters) after the fit
- result : *lmfit.fit\_result Object* container of all information on the fit

3.5.44 fit.voigt\_fit\_2D(x\_scale, y\_scale, data, parameters,  $\lim_{}$ f1,  $\lim_{}$ f2, acqus, N=None, procs=None, utol=(1,1), s1tol=(0,500), s2tol=(0,500), vary utol=(1,1), utol=(0,500), utol=(0,500)

Function that performs the fit of a 2D peak using multiple components. The program reads a parameter matrix, that contains:

u1 /ppm, u2 /ppm, fwhm1 /Hz, fwhm2 /Hz, I /a.u., x\_g

in each row. The number of rows corresponds to the number of components used for the computation of the final signal. The function returns the analogue version of the parameters matrix, but with the optimized values.

## Parameters:

- x\_scale: 1darray ppm\_f2 of the spectrum, full
- y\_scale: 1darray ppm\_f1 of the spectrum, full
- data: *2darray* spectrum, full
- parameters: 1darray or 2darray
  Matrix (# signals, 6). Read main caption.
- lim\_f2: tuple
  Trimming limits for x scale
- lim\_f1: tuple
  Trimming limits for y scale
- acqus: *dict*Dictionary of acquisition parameters.
- N: tuple of ints len(x\_scale). Used only if procs is None
- procs: *dict*Dictionary of processing parameters.
- utol: tuple of floats

  Tolerance for the chemical shifts (utol\_f1, utol\_f2). Values will be set to  $u_1 \pm \text{utol}_{f1}, u_2 \pm \text{utol}_{f2}$ .
- s1tol: tuple of floats
  Range of variations for the fwhm in f1, in Hz
- s2tol: tuple of floats
  Range of variations for the fwhm in f2, in Hz
- vary\_xg: boolChoose if to fix the  $x_q$  value or not
- logfile: str or None
  Path to a file where to write the fit information. If it is None, they will be printed into standard output.

# Returns:

• out\_parameters: 2darray parameters, but with the optimized values.

3.5.45 fit.voigt\_fit\_P2D(S, ppm\_scale, regions, t\_AQ, SFO1, o1p, u\_tol=1, f\_tol=10, vary\_phase=False, vary\_b=False, itermax=10000, file-name='fit')

Performs a lineshape deconvolution fit on a pseudo-2D experiment using a Voigt model. The initial guess must be read from a .ivf file. All components are treated as independent, regardless from the value of the 'group' attribute. The fitting procedure operates iteratively one window at the time. During the fit routine, the peak positions and lineshapes will be varied consistently on all the experiments; only the intensities are allowed to change in a different way.

- S: 2darray Experimental spectrum
- ppm\_scale: 1darray
  PPM scale of the spectrum
- regions: dict Generated by fit.read\_vf\_P2D
- t\_AQ: 1darray
  Acquisition timescale
- SFO1: *float* Nucleus Larmor frequency /MHz
- o1p: float
  Carrier frequency /ppm
- u\_tol: *float*Maximum allowed displacement of the chemical shift from the initial value /ppm
- f\_tol: *float*Maximum allowed displacement of the linewidth from the initial value /ppm
- vary\_phase: bool
  Allow the peaks to change phase
- vary\_b: bool Allow the peaks to change Lorentzian/Gaussian ratio
- itermax: *int*Maximum number of allowed iterations
- filename: strName of the file where the fitted values will be saved. The .fvf extension is added automatically

 $3.5.46 \quad \text{fit.voigt\_fit\_indep}(S, ppm\_scale, regions, t\_AQ, SFO1, o1p, u\_lim=1, \\ f\_lim=10, \ k\_lim=(0, \ 3), \ vary\_phase=False, \ vary\_b=True, \ itermax=10000, \ \text{fit\_tol}=1e-08, \ \text{filename}='\text{fit'}, \ method='leastsq', \ basl\_fit='range' and \ bask of the property of the prop$ 

Performs a lineshape deconvolution fit using a Voigt model. The initial guess must be read from a .ivf file. All components are treated as independent, regardless from the value of the 'group' attribute. The fitting procedure operates iteratively one window at the time.

- S: 1darray Experimental spectrum
- ppm\_scale: 1darray PPM scale of the spectrum
- regions: dict Generated by fit.read\_vf
- t\_AQ: 1darray Acquisition timescale
- SFO1: *float* Nucleus Larmor frequency /MHz
- o1p: float Carrier frequency /ppm
- u\_lim: *float*Maximum allowed displacement of the chemical shift from the initial value /ppm
- f\_lim: float
  Maximum allowed displacement of the linewidth from the initial value /ppm
- k\_lim: float or tuple

  If tuple, minimum and maximum allowed values for k during the fit. If float, maximum displacement from the initial guess
- vary\_phase: bool
  Allow the peaks to change phase
- vary\_b: bool
  Allow the peaks to change Lorentzian/Gaussian ratio
- itermax: *int*Maximum number of allowed iterations
- fit\_tol: float
  Target value to be set for x tol and f tol
- filename: strName of the file where the fitted values will be saved. The .fvf extension is added automatically
- method: str or list of str Method to be used for the optimization. See lmfit for details. There is the option to run multiple optimizations in series.

- basl\_fit: strHow to address the baseline fit. The options are:
  - 'no': Do not use baseline (default)
  - 'fixed': The baseline is computed once and kept fixed during the optimization
  - 'fit': The baseline coefficients enter as fit parameters during the nonlinear optimization
  - 'calc': The baseline coefficients are calculated during the optimization via linear least-squares optimization

## Returns:

• lmfit\_results: list of lmfit.minimizer.MinimizerResult Sequence of the fit results, ordered as the regions dictionary

# 3.5.47 fit.write\_log(input\_file, output\_file, limits, V\_i, C\_i, V\_f, C\_f, result, runtime, test\_res=True, log\_file='fit.log')

Write a log file with all the information of the fit.

- input\_file: str Location and filename of the input file
- output\_file: strLocation and filename of the output file
- limits: *tuple*Delimiters of the spectral region that was fitted. (left, right)
- V\_i: 2darray
  Initial parameters of the fit
- C\_i: 1darray or False
  Coefficients of the starting polynomion used for baseline correction. If False, it was not used.
- V\_f: 2darray Final parameters of the fit
- C\_f: 1darray or False Coefficients of the final polynomion used for baseline correction. If False, it was not used.
- result: *lmfit.FitResult Object*Object returned by lmfit after the fit.
- runtime: datetime.datetime Object Time taken for the fit
- test\_res: bool Choose if to test the residual with the fit.test\_residual function (True) or not (False)
- log\_file: strFilename of the log file to be saved.

## 3.5.48 fit.write par(V, C, limits, filename='i guess.inp')

Write the parameters of the fit, whether they are input or output.

- V : 2darray matrix (# signals, parameters)
- C: 1darray or False
  Coefficients of the polynomion to be used as baseline correction. If the 'baseline' checkbox in the interactive figure panel is not checked, C\_f is False.
- limits: tuple
  Trim limits for the spectrum (left, right).
- $\bullet$  filename: str directory and name of the file to be written

## 3.5.49 fit.write vf(filename, peaks, lims, I, prev=0, header=False, bas c=None

Write a section in a fit report file, which shows the fitting region and the parameters of the peaks to feed into a Voigt lineshape model.

#### Parameters:

• filename: strPath to the file to be written

• peaks: *dict*Dictionary of fit.Peak objects

• lims: tuple (left limit /ppm, right limit /ppm)

• I: *float*Absolute intensity value

• prev: *int*Number of previous peaks already saved. Increases the peak index

• header: bool

If True, adds a '!' starting line to separate fit trials

• bas\_c: None or 1darray
Baseline coefficients

## 3.5.50 fit.write vf P2D(filename, peaks, lims, prev=0)

Write a section in a fit report file, which shows the fitting region and the parameters of the peaks to feed into a Voigt lineshape model.

#### Parameters:

• filename: strPath to the file to be written

• peaks: *list of dict* list of dictionares of fit.Peak objects, one per experiment

• lims: *tuple* (left limit /ppm, right limit /ppm)

• prev: *int*Number of previous peaks already saved. Increases the peak index

## 3.6 SPECTRA package

All the classes in the Spectra module are automatically imported together with klassez itself.

Refer to the examples reported in the *User guide* section to understand how to use them, or use the functions help(), vars(), dir() to get detailed info on how they exactly work.

## 3.6.1 Spectra.Pseudo 2D

class

Subclass of Spectrum\_2D to simulate and handle pseudo-2D experiments. Basically, they share more or less the same attributes, but some methods were adapted in order to suit well with a not-Fourier-transformed indirect dimension.

#### Attributes:

• datadir: *str*Path to the input file/dataset directory

• filename: strBase of the name of the file, without extensions

• fid: 2darray
FID. For simulated data, this must be explicitly set!

• acqus: *dict*Dictionary of acqusition parameters

• ngdic: dict Created only if it is an experimental spectrum. Generated by nmrglue.bruker.read, contains all the information on the spectrometer and on the spectrum.

• procs: *dict*Dictionary of processing parameters

• S: 2darray
Complex spectrum

• rr: 2darray Real part F2, real part F1

• ii: 2darray Imaginary part F2, imaginary part F1

• freq\_f1: 1darray
Indeces of the experiments, works as placeholder

• freq\_f2: 1darray
Frequency scale of the direct dimension, in Hz

• ppm\_f1: *1darray*Indeces of the experiments, works as placeholder

• ppm\_f2: 1darray ppm scale of the direct dimension

• trf1: dict
Projections of the indirect dimension, as 1darrays. Keys: 'ppm\_f2' where they were taken

• trf2: dict

Projections of the direct dimension, as 1darrays. Keys: 'ppm f1' where they were taken

• Trf1: dict

Projections of the indirect dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f2' where they were taken

• Trf2: dict

Projections of the direct dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f1' where they were taken

• integrals: dict

Dictionary where to save the regions and values of the integrals.

• F: fit. Voigt\_Fit\_P2D object Interface for lineshape deconvolution.

#### Methods:

\_\_init\_\_(self, in\_file, pv=False, isexp=True)

Initialize the class.

#### Parameters:

- in\_file: str
  path to file to read, or to the folder of the spectrum
- pv: bool
- True if you want to use pseudo-voigt lineshapes for simulation, False for Voigt
- isexp: bool

True if this is an experimental dataset, False if it is simulated

## add\_noise(self, s\_n=1)

Adds noise to the FID, using the function sim.noisegen.

#### Parameters:

• s\_n: *float*Standard deviation of the noise

## adjph(self, expno=0, p0=None, p1=None, pv=None, update=True)

Adjusts the phases of the spectrum according to the given parameters, or interactively if they are left as default.

- expno: *int*Index of the experiment (python numbering) to use in the interactive panel
- p0: float or None 0-th order phase correction /°
- p1: float or None 1-st order phase correction /°
- pv: float or None 1-st order pivot /ppm
- update: bool
  Choose if to upload the procs dictionary or not

## align(self, lims=None, u\_off=0.5, ref\_idx=0)

Aligns the spectrum to a reference signal in the reference spectrum (default: first one).

#### Parameters:

- lims: tuple or None
  Reference signal region, in ppm. If None, you can select it interactively.
- u\_off: *float*Maximum displacement allowed, in ppm
- ref\_idx: int
  Index of the spectrum to be used as a reference (python numbering)

## basl(self, from procs=False, phase=True)

Apply baseline correction to the whole pseudo-2D by subtracting self.baseline from self.S. Then, self.S is unpacked in self.rr and self.ii.

#### Parameters:

- from \_procs: bool If True, computes the baseline using the polynomion model reading self.procs['basl\_c'] as coefficients
- phase: bool
  Choose if to apply the same phase correction of the spectrum to the baseline. This should be done if the baseline was computed before the phase adjustment!

#### cal(self, offset=None, isHz=False, update=True)

Calibration of the ppm and frequency scales according to a given value, or interactively. In this latter case, a reference peak must be chosen. Calls processing.calibration

• offset: float scale shift F2

• isHz: tuple of bool

True if offset is in frequency units, False if offset is in ppm

 $\bullet$  update: bool

Choose if to update the procs dictionary or not

## calf1(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the indirect dimension according to a given value, or interactively. Calls self.cal on F1 only.

#### Parameters:

• value: float or None scale shift value

• isHz: bool

True if offset is in frequency units, False if offset is in ppm

## calf2(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the direct dimension according to a given value, or interactively. Calls self.cal on F2 only

#### Parameters:

• value: float or None scale shift value

• isHz: bool

True if offset is in frequency units, False if offset is in ppm

#### convdta(self, scaling=1)

Calls processing.convdta

#### eae(self)

Calls processing.EAE to shuffle the data and make a States-like FID. Sets self.eaeflag to 0.

#### integrate(self, which=0, lims=None)

Integrate the spectrum with a dedicated GUI. Calls processing.integral on each experiment, then saves the results in self.integrals. Therefore, the entries of self.integrals are sequences! If lims is not given, calls fit.integrate on the trace to select the regions to integrate.

 $\bullet$  which: int

Experiment index to show in interactive panel

• lims: tuple

Region of the spectrum to integrate (ppm1, ppm2)

## inv process(self)

Performs the inverse processing of the spectrum according to the given parameters. Overwrites the S attribute!! Calls inv xfb.

#### mc(self)

Computes the magnitude of the spectrum on self.S. Then, updates rr, ri, ir, ii.

## mount(self, fids=[], filename=None, newacqus=None)

Replaces the FID of the experiment with a custom one, made by stacking 1D experiments. If the default filename exists (i.e. '<self.filename>.npy'), the function loads it, otherwise calls processing.stack\_fids to create it. The 'fid' attribute is overwritten. The key TD1 of the acqus dictionary is updated to match the first dimension of the new FID.

#### Parameters:

- fids: sequence of 1darray or Spectrum\_1D objects
  FIDs to be stacked. It can be empty if the .npy file already exists.
- filename: str or None
  Path to the filename, without the .npy extension. If it is None, the default filename is used.
- newacqus: dict New acqus dictionary that replaces the actual one. If it is not a dictionary, no actions are performed.

#### pknl(self)

Reverses the effect of the digital filter by applying a first order phase correction. To be called after having processed the data by 'self.process()'

## plot(self, Neg=True, lvl0=0.2, Y label=")

Plots the real part of the spectrum as a 2D contour plot.

- Neg: bool
  Plot (True) or not (False) the negative contours.
- lvl0: float Starting contour value.
- Y\_label: str Custom label for vertical axis.

#### plot md(self, which=None, lims=None)

Plot a number of experiments, superimposed.

#### Parameters:

- which: str or None
  List of experiment indexes, so that eval(which) is meaningful. None plots all of them
- lims: tuple
  Region of the spectrum to show (ppm1, ppm2)

## plot\_stacked(self, which=None, lims=None)

Plot a number of experiments, stacked.

#### Parameters:

- which: str or None List of experiment indexes, so that eval(which) is meaningful. None plots all of them.
- lims: tuple
  Region of the spectrum to show (ppm1, ppm2)

## process(self)

Process only the direct dimension. Calls processing.fp on each transient. The parameters are read from the procs dictionary

#### projf1(self, a, b=None)

Calculates the sum trace of the indirect dimension, from a to b in F2. Store the trace in the dictionary trf1 and as 1D spectrum in Trf1. The key is 'a' or 'a:b' Updates the Trf1[label].freq and Trf1[label].ppm with self.freq\_f1 and self.ppm\_f1 respectively.

#### Parameters:

- a: float ppm F2 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F2.

#### projf2(self, a, b=None)

Calculates the sum trace of the direct dimension, from a to b in F1. Store the trace in the dictionary trf2 and as 1D spectrum in Trf2. The key is 'a' or 'a:b'

- a: *float* ppm F1 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F1.

#### qfil(self, which=None, u=None, s=None)

Gaussian filter to suppress signals. Tries to read self.procs['qfil'], which is { 'u': u, 's': s } Otherwise, these are set interactively by processing.interactive\_qfil and then added to self.procs. Calls processing.qfil

#### Parameters:

- which: int or None Index of the F2 trace to be used for interactive\_qfil. If None, a suitable trace can be selected using misc.select\_traces.
- u: float
  Position /ppm
- s: float
  Width (standard deviation) /ppm

## read\_procs(self, other\_dir=None)

Reads the procs dictionary from a file named 'filename.procs' in the same directory of the input file.

#### Parameters:

• other\_dir: str or None
Different location for the procs dictionary to look into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### Returns:

• procs: *dict*Dictionary of processing parameters

## rpbc(self, ref exp=0, \*\*rpbc kws)

Computes the phase angles and the baseline using processing.rpbc on a reference spectrum taken from self.S. Then applies the phase correction and subtracts the baseline, automatically, to all experiments of the pseudo-2D. The procs dictionary is then updated and saved. The polynomial baseline is computed according to the given coefficients and stored in self.baseline

- ref\_exp: int
  Index of the reference experiment on which to apply the algorithm
- rpbc\_kws: keyworded arguments See processing.RPBC for details.

## scan(self, ns=1, s n=1)

Simulates the acquisition of ns scans, by adding a different realization of noise at each iteration. The function is supposed to start with the FID without noise at all. If not, the results will be biased.

#### Parameters:

- ns: *int*Number of scans to accumulate
- s\_n: *float* Standard deviation of the noise

## to wav(self, filename=None, cutoff=None, rate=44100)

Converts the FID in an audio file by using misc.data2wav.

#### Parameters:

- $\bullet$  filename: strPath where to save the file. If None, self.filename is used
- cutoff: *float* Clipping limits for the FID
- rate: *int* Sampling rate in samples/sec

## write\_acqus(self, other\_dir=None)

Write the acqus dictionary in a file named 'filename.acqus'. Calls misc.write acqus 1D

#### Parameters:

• other\_dir: str or None
Different location for the acqus dictionary to write into. If None, self.datadir is used instead.

#### write integrals(self, filename='integrals.dat')

Write the integrals in a file named filename.

• filename: str name of the file where to write the integrals.

## write procs(self, other dir=None)

Writes the actual procs dictionary in a file named 'filename.procs' in the same directory of the input file.

#### Parameters:

• other\_dir: str or None
Different location for the procs dictionary to write into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

## write ser(ser, acqus, path=None)

Writes a real/complex array in binary format. Calls misc.write\_ser. Be sure that acqus contains the BYTORDA and DTYPA keys. See misc.write\_ser to understand the meaning of these values.

#### Parameters:

- ser: *ndarray*Array that you want to convert in binary format.
- acqus: dict
  Dictionary of acquisition parameters. It must contain BYTORDA and DTYPA.
- path: *str*Path where to save the binary file.

#### xf1(self)

Process only the indirect dimension. Transposes the spectrum in hypermode or normally if FnMODE!= QF, then calls for processing.fp using self.procs[keys][0], finally transposes it back. The result is stored in self.S, then self.rr and self.ii are written. freq\_f1 and ppm\_f1 are assigned with the indexes of the transients.

#### xf2(self)

Process only the direct dimension. Calls processing.fp using procs[keys][1] The result is stored in self.S, then self.rr and self.ii are written. freq\_f1 and ppm\_f1 are assigned with the indexes of the transients.

## 3.6.2 Spectra. Spectrum 1D

class

Class: 1D NMR spectrum

#### Attributes:

• datadir: *str*Path to the input file/dataset directory

• filename: strBase of the name of the file, without extensions

• fid: *1darray* FID

• acqus: *dict*Dictionary of acqusition parameters

• ngdic: dict Created only if it is an experimental spectrum. Generated by nmrglue.bruker.read, contains all the information on the spectrometer and on the spectrum.

• procs: *dict*Dictionary of processing parameters

• S: *1darray*Complex spectrum

• r: *1darray*Real part of the spectrum

• i: *1darray*Imaginary part of the spectrum

• freq: *1darray*Frequency scale of the spectrum, in Hz

• ppm: *1darray* ppm scale of the spectrum

• F: fit. Voigt\_Fit object Used for deconvolution. See fit. Voigt fit.

• baseline: *1darray*Baseline of the spectrum.

• integrals: *dict*Dictionary where to save the regions and values of the integrals.

#### Methods:

```
__init__(self, in_file, pv=False, isexp=True, spect='bruker')
```

Initialize the class. Simulation of the dataset (i.e. isexp=False) employs sim.sim\_1D.

- in\_file: str
  path to file to read, or to the folder of the spectrum
- pv: bool

  True if you want to use pseudo-voigt lineshapes for simulation, False for Voigt
- isexp: bool

  True if this is an experimental dataset, False if it is simulated
- spect: strData file format. Allowed: 'bruker', 'varian', 'magritek', 'oxford', 'jeol'

## acme(self, \*\*method kws)

Automatic phase correction based on entropy minimization It calculates the phase angles using the algorithm specified in method, then calls self.adjph with those values.

#### Parameters:

• method\_kws: keyworded arguments
Additional parameters for the chosen method.

## add noise(self, s n=1)

Adds noise to the FID, using the function sim.noisegen.

#### Parameters:

• s\_n: *float* Standard deviation of the noise

#### adjph(self, p0=None, p1=None, pv=None, update=True)

Adjusts the phases of the spectrum according to the given parameters, or interactively if they are left as default. Calls for processing.ps

- p0: float or None 0-th order phase correction /°
- p1: float or None 1-st order phase correction /°
- pv: float or None 1-st order pivot /ppm
- update: bool Choose if you want to update the procs dictionary or not

## baseline correction(self, basl file='spectrum.basl', winlim=None)

Correct the baseline of the spectrum, according to a pre-existing file or interactively. Calls processing.baseline correction or processing.load baseline

#### Parameters:

- basl\_file: str
  Path to the baseline file. If it already exists, the baseline will be built according to this file;
  otherwise this will be the destination file of the baseline.
- winlim: tuple or None Limits of the baseline. If it is None, it will be interactively set. If basl\_file exists, it will be read from there. Else, (ppm1, ppm2).

## basl(self, from procs=False, phase=True)

Apply the baseline correction by subtracting self.baseline from self.S. Then, self.S is unpacked in self.r and self.i

#### Parameters:

- from \_procs: bool

  If True, computes the baseline using the polynomion model reading self.procs['basl\_c'] as coefficients
- phase: bool
  Choose if to apply the same phase correction of the spectrum to the baseline. This should be done if the baseline was computed before the phase adjustment!

#### blp(self, pred=8, order=8)

Call processing.blp on self.fid for the application of backward linear prediction to the data. Important for Oxford benchtop data, where you have to predict 8 points to have a usable spectrum.

#### Parameters:

- pred: *int*Number of points to be predicted
- order: *int*Number of coefficients to be used for the prediction
- N: int
  Number of FID points to be used for calculation; used to decrease computation time

#### cal(self, offset=None, isHz=False, update=True)

Calibrates the ppm and frequency scale according to a given value, or interactively. Calls processing.calibration

- offset: float or None scale shift value
- isHz: bool

True if offset is in frequency units, False if offset is in ppm

• update: bool

Choose if to update the procs dictionary or not

#### convdta(self, scaling=1)

Call processing.convdta using self.acqus['GRPDLY']

## integrate(self, lims=None)

Integrate the spectrum with a dedicated GUI. Calls fit.integrate and writes in self.integrals with keys [ppm1:ppm2]

#### Parameters:

• lims: tuple
Integrates from lims[0] to lims[1]. If it is None, calls for interactive integration.

## inv\_process(self)

Performs the inverse processing of the spectrum according to the given parameters. Overwrites the S attribute!! Calls processing.inv\_fp

#### mc(self)

Calculates the magnitude of the spectrum and overwrites self.S, self.r, self.i

## pknl(self)

Reverses the effect of the digital filter by applying a first order phase correction. To be called after having processed the data by 'self.process()'

#### plot(self, name=None, ext='png', dpi=600)

Plots the real part of the spectrum.

#### Parameters:

- $\bullet$  name: str
  - Filename for the figure. If None, it is shown instead.
- $\bullet$  ext: str

Format of the image

• dpi: int

Resolution of the image in dots per inches

#### process(self, interactive=False)

Performs the processing of the FID. The parameters are read from self.procs. Calls processing.interactive\_fp or processing.fp using self.acqus and self.procs Writes the result is self.S, then unpacks it in self.r and self.i Calculates frequency and ppm scales. Also initializes self.F with fit.Voigt Fit class using the current parameters

#### **Parameters:**

• interactive: bool

True if you want to open the interactive panel, False to read the parameters from self.procs.

#### qfil(self, u=None, s=None)

Gaussian filter to suppress signals. Tries to read self.procs['qfil'], which is { 'u': u, 's': s } Otherwise, these are set interactively by processing.interactive\_qfil and then added to self.procs. Calls processing.qfil

#### Parameters:

- u: *float*Position of the filter /ppm
- s: float
  Width (standard deviation) of the filter /ppm

## read\_procs(self, other\_dir=None)

Reads the procs dictionary from a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to look into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### **Returns:**

• procs: dict
Dictionary of processing parameters

## rpbc(self, \*\*rpbc\_kws)

Computes the phase angles and the baseline using processing.RPBC on self.S. Then applies the phase correction and subtracts the baseline, automatically. The procs dictionary is then updated and saved. The polynomial baseline is computed according to the given coefficients and stored in self.baseline

• rpbc\_kws: keyworded arguments See processing.RPBC for details.

## scan(self, ns=1, s n=1)

Simulates the acquisition of ns scans, by adding a different realization of noise at each iteration. The function is supposed to start with the FID without noise at all. If not, the results will be biased.

#### Parameters:

- ns: *int*Number of scans to accumulate
- s\_n: *float* Standard deviation of the noise

## to vf(self, filename=None, Hs=None, fvf=True)

Transform a simulated spectrum in a .ivf or .fvf file to be used in a deconvolution procedure. To do this, it reads the peak parameters saved in acqus. The number of signals is determined by acqus['amplitudes']. Multiplets are splitted according to their Js, and saved as components.

#### Parameters:

- filename:  $str\ or\ None$ Path to the filename to be saved, without extension. If None, self.filename is used
- Hs: int or None Number of nuclei the spectrum integrates for. If None, the sum of the amplitudes is used.
- fvf: bool

  If True, adds the '.fvf' extension to the filename, if False, adds '.ivf'

## to wav(self, filename=None, cutoff=None, rate=44100)

Converts the FID in an audio file by using misc.data2wav.

- filename: strPath where to save the file. If None, self.filename is used
- cutoff: *float* Clipping limits for the FID
- rate: *int* Sampling rate in samples/sec

## write acqus(self, other dir=None)

Write the acqus dictionary in a file named 'filename.acqus'. Calls misc.write acqus 1D

#### **Parameters:**

• other\_dir: str or None
Different location for the acqus dictionary to write into. If None, self.datadir is used instead.

## write integrals(self, other dir=None)

Write the integrals in a file named '{self.filename}.int'.

#### Parameters:

• other\_dir: str or None

Different location for the integrals file to write into. If None, self.datadir is used instead.

## write\_procs(self, other\_dir=None)

Writes the actual procs dictionary in a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to write into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

## write ser(ser, acqus, path=None)

Writes a real/complex array in binary format. Calls misc.write\_ser. Be sure that acqus contains the BYTORDA and DTYPA keys. See misc.write\_ser to understand the meaning of these values.

- ser: *ndarray*Array that you want to convert in binary format.
- acqus: dict Dictionary of acquisition parameters. It must contain BYTORDA and DTYPA.
- path: strPath where to save the binary file.

## 3.6.3 Spectra.Spectrum 2D

class

Class: 2D NMR spectrum

#### Attributes:

 $\bullet$  datadir: str

Path to the input file/dataset directory

 $\bullet$  filename: str

Base of the name of the file, without extensions

• fid: 2darray

FID

• acqus: dict

Dictionary of acqusition parameters

• ngdic: dict

Created only if it is an experimental spectrum. Generated by nmrglue.bruker.read, contains all the information on the spectrometer and on the spectrum.

• procs: dict

Dictionary of processing parameters

• eaeflag: int

If FnMODE is Echo-Antiecho, keeps track of the manipulation of the data so to not repeat the same process twice

• S: 2darray

Complex (or hypercomplex, depending on FnMODE) spectrum

• rr: 2darray

Real part F2, real part F1

• ii: 2darray

Imaginary part F2, imaginary part F1

• ir: 2darray

Real part F2, imaginary part F1. Only exist if F1 is acquired in phase-sensitive mode

• ri: 2darray

Imaginary part F2, real part F1. Only exist if F1 is acquired in phase-sensitive mode

• freq f1: 1darray

Frequency scale of the indirect dimension, in Hz

• freq f2: 1darray

Frequency scale of the direct dimension, in Hz

• ppm f1: 1darray

ppm scale of the indirect dimension

• ppm f2: 1darray

ppm scale of the direct dimension

trf1: dict

Projections of the indirect dimension, as 1darrays. Keys: 'ppm\_f2' where they were taken

- trf2: dict
  Projections of the direct dimension, as 1darrays. Keys: 'ppm f1' where they were taken
- Trf1: dict Projections of the indirect dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f2' where they were taken
- Trf2: dict Projections of the direct dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f1' where they were taken
- integrals: *dict*Dictionary where to save the regions and values of the integrals.

#### Methods:

\_\_init\_\_(self, in\_file, pv=False, isexp=True, is\_pseudo=False)

Initialize the class.

#### **Parameters:**

- in\_file: str
  path to file to read, or to the folder of the spectrum
- pv: bool

  True if you want to use pseudo-voigt lineshapes for simulation, False for Voigt
- isexp: bool

  True if this is an experimental dataset, False if it is simulated
- is\_pseudo: bool True if it is a pseudo-2D. Legacy option

## add\_noise(self, s\_n=1)

Adds noise to the FID, using the function sim.noisegen.

#### Parameters:

• s\_n: *float* Standard deviation of the noise

## 

Adjusts the phases of the spectrum according to the given parameters, or interactively if they are left as default. The non-interactive workflow is to apply processing.ps on F2, transpose according to FnMODE, apply processing.ps on F1, transpose back. If FnMODE is 'No', the phase correction is applied only on F2, as it should be done in a pseudo-2D experiment. Once self.S was updated and unpacked, the phase values are added to the procs dictionary to keep track of multiple phase adjustments.

- p01: float or None 0-th order phase correction /° of the indirect dimension
- p11: float or None 1-st order phase correction /° of the indirect dimension
- pv1: float or None
  1-st order pivot /ppm of the indirect dimension
- p02: float or None 0-th order phase correction /° of the direct dimension
- p12: float or None 1-st order phase correction /° of the direct dimension
- pv2: float or None
  1-st order pivot /ppm of the direct dimension
- update: bool
  Choose if to update the procs dictionary or not

## cal(self, offset=[None, None], isHz=False, update=True)

Calibration of the ppm and frequency scales according to a given value, or interactively. In this latter case, a reference peak must be chosen. Calls processing.calibration

#### Parameters:

- offset: tuple (scale shift F1, scale shift F2)
- isHz: tuple of bool

  True if offset is in frequency units, False if offset is in ppm
- update: bool
  Choose if to update the procs dictionary or not

#### calf1(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the indirect dimension according to a given value, or interactively. Calls self.cal on F1 only.

- value: float or None scale shift value
- isHz: bool

  True if offset is in frequency units, False if offset is in ppm

#### calf2(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the direct dimension according to a given value, or interactively. Calls self.cal on F2 only

#### Parameters:

- value: float or None scale shift value
- isHz: bool

True if offset is in frequency units, False if offset is in ppm

#### convdta(self, scaling=1)

Calls processing.convdta to compensate for the group delay. It does not always work, depends on TopSpin version and planets alignment.

#### Parameters:

• scaling: *float*Scaling factor for processing convolta.

#### eae(self)

Calls processing. EAE to shuffle the data and make a States-like FID. Sets self-eaeflag to 0.

## integrate(self, \*\*kwargs)

Integrates the spectrum with a dedicated GUI. Calls fit.integrate\_2D

#### Parameters:

• kwargs: keyworded arguments
Additional parameters for fit.integrate\_2D

## inv process(self)

Performs the inverse processing of the spectrum according to the given parameters. Overwrites the S attribute!! Calls inv\_xfb.

#### mc(self)

Computes the magnitude of the spectrum on self.S. Then, updates rr, ri, ir, ii.

#### pknl(self)

Reverses the effect of the digital filter by applying a first order phase correction. To be called after having processed the data by 'self.process()'

#### plot(self, Neg=True, lvl0=0.2)

Plots the real part of the spectrum. Use the mouse scroll to adjust the contour starting level.

#### Parameters:

- Neg: bool
  Plot (True) or not (False) the negative contours.
- lvl0: float
  Starting contour value with respect to the maximum of the spectrum

## process(self, interactive=False, \*\*int kwargs)

Performs the full processing of the FID on both dimensions. The parameters are read from self.procs. If FnMODE is Echo-Antiecho and you did not call self.eae before, the FID is converted to States with processing.EAE before to start. If interactive is True, calls processing.interactive\_xfb with int\_kwargs, else calls processing.xfb. The complex/hypercomplex spectrum is stored in self.S, then unpacked into self.rr, self.ri, self.ir, self.ii. If FnMODE is Echo-Antiecho, a phase correction of -90 degrees is applied on the indirect dimension.

#### Parameters:

- interactive: bool

  True if you want to open the interactive panel, False to read the parameters from self.procs.
- int\_kwargs: keyworded arguments
  Additional parameters for processing.interactive\_xfb, if interactive=True.

#### projf1(self, a, b=None)

Calculates the sum trace of the indirect dimension, from a ppm to b ppm in F2. Store the trace in the dictionary trf1 and as 1D spectrum in Trf1. The key is 'a' or 'a:b' Calls misc.get\_trace on self.rr with column=True

#### Parameters:

- a: *float* ppm F2 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F2.

#### projf2(self, a, b=None)

Calculates the sum trace of the direct dimension, from a ppm to b ppm in F1. Store the trace in the dictionary trf2 and as 1D spectrum in Trf2. The key is 'a' or 'a:b' Calls misc.get\_trace on self.rr with column=False

- a: *float* ppm F1 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F1.

## qfil(self, which=None, u=None, s=None)

Gaussian filter to suppress signals. Tries to read self.procs['qfil'], which is { 'u': u, 's': s } Otherwise, these are set interactively by processing.interactive\_qfil and then added to self.procs. Calls processing.qfil

#### **Parameters:**

- which: int or None Index of the F2 trace to be used for interactive\_qfil. If None, a suitable trace can be selected using misc.select\_traces.
- u: float
  Position /ppm
- s: float
  Width (standard deviation) /ppm

## read procs(self, other dir=None)

Reads the procs dictionary from a file named 'filename.procs' in the same directory of the input file.

## Parameters:

• other\_dir: str or None
Different location for the procs dictionary to look into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### **Returns:**

• procs: *dict*Dictionary of processing parameters

#### scan(self, ns=1, s n=1)

Simulates the acquisition of ns scans, by adding a different realization of noise at each iteration. The function is supposed to start with the FID without noise at all. If not, the results will be biased.

- ns: *int* Number of scans to accumulate
- s\_n: *float*Standard deviation of the noise

## to wav(self, filename=None, cutoff=None, rate=44100)

Converts the FID in an audio file by using misc.data2wav.

#### Parameters:

- $\bullet$  filename: strPath where to save the file. If None, self.filename is used
- cutoff: *float* Clipping limits for the FID
- rate: *int* Sampling rate in samples/sec

#### write acqus(self, other dir=None)

Write the acqus dictionary in a file named 'filename.acqus'. Calls misc.write\_acqus\_1D

#### **Parameters:**

• other\_dir: str or None
Different location for the acqus dictionary to write into. If None, self.datadir is used instead.

## write\_integrals(self, other\_dir=None)

Write the integrals in a file named '{self.filename}.int'.

#### Parameters:

• other\_dir: str or None Different location for the integrals file to write into. If None, self.datadir is used instead.

## write\_procs(self, other\_dir=None)

Writes the actual procs dictionary in a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to write into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### write ser(ser, acqus, path=None)

Writes a real/complex array in binary format. Calls misc.write\_ser. Be sure that acqus contains the BYTORDA and DTYPA keys. See misc.write\_ser to understand the meaning of these values.

• ser: *ndarray*Array that you want to convert in binary format.

• acqus: dict Dictionary of acquisition parameters. It must contain BYTORDA and DTYPA.

• path: strPath where to save the binary file.

## xf1(self)

Process only the indirect dimension. Transposes the spectrum in hypermode or normally if FnMODE!= QF, then calls for processing.fp using self.procs[keys][0], finally transposes it back. The result is stored in self.S, then self.rr and self.ii are written. freq\_fl and ppm\_fl are assigned with the indexes of the transients.

## xf2(self)

Process only the direct dimension. Calls processing.fp using procs[keys][1] The result is stored in self.S, then self.rr and self.ii are written. freq\_f1 and ppm\_f1 are assigned with the indexes of the transients.

## 3.6.4 Spectra.pSpectrum 1D

class

Subclass of Spectrum\_1D that allows to handle processed 1D NMR spectra. Useful when dealing with traces of 2D spectra. Shares the same attributes with Spectrum 1D.

#### Attributes:

 $\bullet$  datadir: str

Path to the input file/dataset directory

• filename: str

Base of the name of the file, without extensions

• acqus: dict

Dictionary of acqusition parameters

• ngdic: dict

Created only if it is an experimental spectrum. Generated by nmrglue.bruker.read, contains all the information on the spectrometer and on the spectrum.

• procs: dict

Dictionary of processing parameters

• S: 1darray

Complex spectrum

• r: 1darray

Real part of the spectrum

• i: 1darray

Imaginary part of the spectrum

• freq: 1darray

Frequency scale of the spectrum, in Hz

• ppm: 1darray

ppm scale of the spectrum

• F: fit. Voigt Fit object

Used for deconvolution. See fit. Voigt fit.

• baseline: 1darray

Baseline of the spectrum.

• integrals: dict

Dictionary where to save the regions and values of the integrals.

#### Methods:

init \_\_(self, in\_file, acqus=None, procs=None, istrace=False, filename='T')

Initialize the class.

- in\_file: str or 1darray

  If istrace is True, in file is the NMR spectrum. Else, it is the directory of the processed data.
- acqus: dict or None

  If istrace is True, you must supply the associated 'acqus' dictionary. Else, it is not necessary as it is read from the input directory
- procs: dict or None
  You can pass the dictionary of processing parameters, if you want. Otherwise, it is initialized with standard values.
- istrace: bool

  Declare the object as trace extracted from a 2D (True) or as true experimental spectrum (False)
- $\bullet$  filename: strIf istrace is True, this will be the filename of self.acqus and self.procs

## acme(self, \*\*method\_kws)

Automatic phase correction based on entropy minimization It calculates the phase angles using the algorithm specified in method, then calls self-adjph with those values.

#### Parameters:

• method\_kws: keyworded arguments
Additional parameters for the chosen method.

## add\_noise(self, s\_n=1)

Adds noise to the FID, using the function sim.noisegen.

#### Parameters:

• s\_n: *float* Standard deviation of the noise

#### adjph(self, p0=None, p1=None, pv=None, update=True)

Adjusts the phases of the spectrum according to the given parameters, or interactively if they are left as default. Calls for processing.ps

- p0: float or None 0-th order phase correction /°
- p1: float or None 1-st order phase correction /°
- pv: float or None 1-st order pivot /ppm
- update: bool
  Choose if you want to update the procs dictionary or not

## baseline correction(self, basl file='spectrum.basl', winlim=None)

Correct the baseline of the spectrum, according to a pre-existing file or interactively. Calls processing.baseline correction or processing.load baseline

#### Parameters:

- basl\_file: str
  Path to the baseline file. If it already exists, the baseline will be built according to this file; otherwise this will be the destination file of the baseline.
- winlim: tuple or None Limits of the baseline. If it is None, it will be interactively set. If basl\_file exists, it will be read from there. Else, (ppm1, ppm2).

## basl(self, from procs=False, phase=True)

Apply the baseline correction by subtracting self.baseline from self.S. Then, self.S is unpacked in self.r and self.i

#### Parameters:

- from procs: bool

  If True, computes the baseline using the polynomion model reading self.procs['basl\_c'] as coefficients
- phase: bool
  Choose if to apply the same phase correction of the spectrum to the baseline. This should be done if the baseline was computed before the phase adjustment!

#### blp(self, pred=8, order=8)

Call processing.blp on self.fid for the application of backward linear prediction to the data. Important for Oxford benchtop data, where you have to predict 8 points to have a usable spectrum.

#### Parameters:

- pred: *int*Number of points to be predicted
- order: *int*Number of coefficients to be used for the prediction
- N: int
  Number of FID points to be used for calculation; used to decrease computation time

#### cal(self, offset=None, isHz=False, update=True)

Calibrates the ppm and frequency scale according to a given value, or interactively. Calls processing.calibration

• offset: float or None scale shift value

• isHz: bool

True if offset is in frequency units, False if offset is in ppm

• update: bool

Choose if to update the procs dictionary or not

#### convdta(self, scaling=1)

Call processing.convdta using self.acqus['GRPDLY']

## integrate(self, lims=None)

Integrate the spectrum with a dedicated GUI. Calls fit.integrate and writes in self.integrals with keys [ppm1:ppm2]

#### Parameters:

• lims: tuple
Integrates from lims[0] to lims[1]. If it is None, calls for interactive integration.

## inv process(self)

Performs the inverse processing of the spectrum according to the given parameters. Overwrites the S attribute!! Calls processing.inv\_fp

#### mc(self)

Calculates the magnitude of the spectrum and overwrites self.S, self.r, self.i

## pknl(self)

Reverses the effect of the digital filter by applying a first order phase correction. To be called after having processed the data by 'self.process()'

#### plot(self, name=None, ext='png', dpi=600)

Plots the real part of the spectrum.

#### Parameters:

 $\bullet$  name: str

Filename for the figure. If None, it is shown instead.

 $\bullet$  ext: str

Format of the image

• dpi: *int* 

Resolution of the image in dots per inches

#### process(self, interactive=False)

Performs the processing of the FID. The parameters are read from self.procs. Calls processing.interactive\_fp or processing.fp using self.acqus and self.procs Writes the result is self.S, then unpacks it in self.r and self.i Calculates frequency and ppm scales. Also initializes self.F with fit.Voigt\_Fit class using the current parameters

#### **Parameters:**

• interactive: bool

True if you want to open the interactive panel, False to read the parameters from self.procs.

#### qfil(self, u=None, s=None)

Gaussian filter to suppress signals. Tries to read self.procs['qfil'], which is { 'u': u, 's': s } Otherwise, these are set interactively by processing.interactive\_qfil and then added to self.procs. Calls processing.qfil

#### Parameters:

- u: *float*Position of the filter /ppm
- s: float
  Width (standard deviation) of the filter /ppm

#### read procs(self, other dir=None)

Reads the procs dictionary from a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to look into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### **Returns:**

• procs: dict
Dictionary of processing parameters

## rpbc(self, \*\*rpbc\_kws)

Computes the phase angles and the baseline using processing.RPBC on self.S. Then applies the phase correction and subtracts the baseline, automatically. The procs dictionary is then updated and saved. The polynomial baseline is computed according to the given coefficients and stored in self.baseline

• rpbc\_kws: keyworded arguments See processing.RPBC for details.

## scan(self, ns=1, s n=1)

Simulates the acquisition of ns scans, by adding a different realization of noise at each iteration. The function is supposed to start with the FID without noise at all. If not, the results will be biased.

#### Parameters:

- ns: *int*Number of scans to accumulate
- s\_n: *float* Standard deviation of the noise

## to vf(self, filename=None, Hs=None, fvf=True)

Transform a simulated spectrum in a .ivf or .fvf file to be used in a deconvolution procedure. To do this, it reads the peak parameters saved in acqus. The number of signals is determined by acqus['amplitudes']. Multiplets are splitted according to their Js, and saved as components.

#### Parameters:

- filename:  $str\ or\ None$ Path to the filename to be saved, without extension. If None, self.filename is used
- Hs: int or None Number of nuclei the spectrum integrates for. If None, the sum of the amplitudes is used.
- fvf: bool

  If True, adds the '.fvf' extension to the filename, if False, adds '.ivf'

## to\_wav(self, filename=None, cutoff=None, rate=44100)

Converts the FID in an audio file by using misc.data2wav.

- filename: strPath where to save the file. If None, self.filename is used
- cutoff: *float* Clipping limits for the FID
- $\bullet$  rate: int Sampling rate in samples/sec

## write acqus(self, other dir=None)

Write the acqus dictionary in a file named 'filename.acqus'. Calls misc.write acqus 1D

#### **Parameters:**

• other\_dir: str or None
Different location for the acqus dictionary to write into. If None, self.datadir is used instead.

## write integrals(self, other dir=None)

Write the integrals in a file named '{self.filename}.int'.

#### Parameters:

• other\_dir: str or None

Different location for the integrals file to write into. If None, self.datadir is used instead.

## write\_procs(self, other\_dir=None)

Writes the actual procs dictionary in a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to write into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

## write ser(ser, acqus, path=None)

Writes a real/complex array in binary format. Calls misc.write\_ser. Be sure that acqus contains the BYTORDA and DTYPA keys. See misc.write\_ser to understand the meaning of these values.

- ser: *ndarray*Array that you want to convert in binary format.
- acqus: dict Dictionary of acquisition parameters. It must contain BYTORDA and DTYPA.
- path: strPath where to save the binary file.

## 3.6.5 Spectra.pSpectrum 2D

class

Subclass of Spectrum\_2D that allows to handle processed 2D NMR spectra. Reads the processed spectrum from Bruker.

#### Attributes:

 $\bullet$  datadir: str

Path to the input file/dataset directory

 $\bullet$  filename: str

Base of the name of the file, without extensions

• acqus: dict

Dictionary of acqusition parameters

• ngdic: dict

Generated by nmrglue.bruker.read, contains all the information on the spectrometer and on the spectrum.

• procs: dict

Dictionary of processing parameters

• S: 2darray

Complex (or hypercomplex, depending on FnMODE) spectrum

• rr: 2darray

Real part F2, real part F1

• ii: 2darray

Imaginary part F2, imaginary part F1

• ir: 2darray

Real part F2, imaginary part F1. Only exist if F1 is acquired in phase-sensitive mode

• ri: 2darray

Imaginary part F2, real part F1. Only exist if F1 is acquired in phase-sensitive mode

• freq f1: 1darray

Frequency scale of the indirect dimension, in Hz

• freq f2: 1darray

Frequency scale of the direct dimension, in Hz

• ppm f1: 1darray

ppm scale of the indirect dimension

• ppm f2: 1darray

ppm scale of the direct dimension

• trf1· dict

Projections of the indirect dimension, as 1darrays. Keys: 'ppm f2' where they were taken

• trf2: dict

Projections of the direct dimension, as 1darrays. Keys: 'ppm f1' where they were taken

• Trf1: *dict* 

Projections of the indirect dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f2' where they were taken

• Trf2: dict

Projections of the direct dimension, as pSpectrum\_1D objects. Keys: 'ppm\_f1' where they were taken

 $\bullet$  integrals: dict

Dictionary where to save the regions and values of the integrals.

#### Methods:

$$init$$
 (self, in file)

Initialize the class.

#### **Parameters:**

• in\_file: str
Path to the spectrum. Here, the 'pdata/#' folder must be specified.

## add\_noise(self, s\_n=1)

Adds noise to the FID, using the function sim.noisegen.

#### **Parameters:**

• s\_n: *float*Standard deviation of the noise

## 

Adjusts the phases of the spectrum according to the given parameters, or interactively if they are left as default. The non-interactive workflow is to apply processing.ps on F2, transpose according to FnMODE, apply processing.ps on F1, transpose back. If FnMODE is 'No', the phase correction is applied only on F2, as it should be done in a pseudo-2D experiment. Once self.S was updated and unpacked, the phase values are added to the procs dictionary to keep track of multiple phase adjustments.

- p01: float or None 0-th order phase correction /° of the indirect dimension
- p11: float or None 1-st order phase correction /° of the indirect dimension
- pv1: float or None
  1-st order pivot /ppm of the indirect dimension

- p02: float or None 0-th order phase correction /° of the direct dimension
- p12: float or None 1-st order phase correction /° of the direct dimension
- pv2: float or None 1-st order pivot /ppm of the direct dimension
- update: *bool*Choose if to update the procs dictionary or not

## cal(self, offset=[None, None], isHz=False, update=True)

Calibration of the ppm and frequency scales according to a given value, or interactively. In this latter case, a reference peak must be chosen. Calls processing.calibration

#### Parameters:

- offset: tuple (scale shift F1, scale shift F2)
- isHz: tuple of bool

  True if offset is in frequency units, False if offset is in ppm
- update: bool Choose if to update the procs dictionary or not

#### calf1(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the indirect dimension according to a given value, or interactively. Calls self.cal on F1 only.

#### Parameters:

- value: float or None scale shift value
- isHz: bool

  True if offset is in frequency units, False if offset is in ppm

#### calf2(self, value=None, isHz=False)

Calibrates the ppm and frequency scale of the direct dimension according to a given value, or interactively. Calls self.cal on F2 only

- value: float or None scale shift value
- isHz: bool

  True if offset is in frequency units, False if offset is in ppm

#### convdta(self, scaling=1)

Calls processing.convdta to compensate for the group delay. It does not always work, depends on TopSpin version and planets alignment.

#### Parameters:

• scaling: *float*Scaling factor for processing convolta.

## eae(self)

Calls processing. EAE to shuffle the data and make a States-like FID. Sets self.eaeflag to 0.

## integrate(self, \*\*kwargs)

Integrates the spectrum with a dedicated GUI. Calls fit.integrate\_2D

#### Parameters:

• kwargs: keyworded arguments
Additional parameters for fit.integrate 2D

## inv process(self)

Performs the inverse processing of the spectrum according to the given parameters. Overwrites the S attribute!! Calls inv\_xfb.

#### mc(self)

Computes the magnitude of the spectrum on self.S. Then, updates rr, ri, ir, ii.

## pknl(self)

Reverses the effect of the digital filter by applying a first order phase correction. To be called after having processed the data by 'self.process()'

#### plot(self, Neg=True, lvl0=0.2)

Plots the real part of the spectrum. Use the mouse scroll to adjust the contour starting level.

- Neg: bool
  Plot (True) or not (False) the negative contours.
- lvl0: float
  Starting contour value with respect to the maximum of the spectrum

## process(self, interactive=False, \*\*int kwargs)

Performs the full processing of the FID on both dimensions. The parameters are read from self.procs. If FnMODE is Echo-Antiecho and you did not call self.eae before, the FID is converted to States with processing.EAE before to start. If interactive is True, calls processing.interactive\_xfb with int\_kwargs, else calls processing.xfb. The complex/hypercomplex spectrum is stored in self.S, then unpacked into self.rr, self.ri, self.ir, self.ii. If FnMODE is Echo-Antiecho, a phase correction of -90 degrees is applied on the indirect dimension.

#### Parameters:

- interactive: bool

  True if you want to open the interactive panel, False to read the parameters from self.procs.
- int\_kwargs: keyworded arguments
  Additional parameters for processing.interactive\_xfb, if interactive=True.

## projf1(self, a, b=None)

Calculates the sum trace of the indirect dimension, from a ppm to b ppm in F2. Store the trace in the dictionary trf1 and as 1D spectrum in Trf1. The key is 'a' or 'a:b' Calls misc.get\_trace on self.rr with column=True

#### Parameters:

- a: float ppm F2 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F2.

#### projf2(self, a, b=None)

Calculates the sum trace of the direct dimension, from a ppm to b ppm in F1. Store the trace in the dictionary trf2 and as 1D spectrum in Trf2. The key is 'a' or 'a:b' Calls misc.get\_trace on self.rr with column=False

#### Parameters:

- a: *float* ppm F1 value where to extract the trace.
- b: float or None.

  If it is None, extract the trace in a. Else, sum from a to b in F1.

#### qfil(self, which=None, u=None, s=None)

Gaussian filter to suppress signals. Tries to read self.procs['qfil'], which is { 'u': u, 's': s } Otherwise, these are set interactively by processing.interactive\_qfil and then added to self.procs. Calls processing.qfil



- which: int or None Index of the F2 trace to be used for interactive\_qfil. If None, a suitable trace can be selected using misc.select traces.
- u: float
  Position /ppm
- s: float
  Width (standard deviation) /ppm

## read procs(self, other dir=None)

Reads the procs dictionary from a file named 'filename.procs' in the same directory of the input file.

#### **Parameters:**

• other\_dir: str or None
Different location for the procs dictionary to look into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### Returns:

• procs: *dict*Dictionary of processing parameters

## scan(self, ns=1, s n=1)

Simulates the acquisition of ns scans, by adding a different realization of noise at each iteration. The function is supposed to start with the FID without noise at all. If not, the results will be biased.

#### Parameters:

- ns: *int*Number of scans to accumulate
- s\_n: *float* Standard deviation of the noise

## $to\_wav(self, filename{=}None, \, cutoff{=}None, \, rate{=}44100)$

Converts the FID in an audio file by using misc.data2wav.

- filename: strPath where to save the file. If None, self.filename is used
- cutoff: *float* Clipping limits for the FID
- rate: *int* Sampling rate in samples/sec

## write acqus(self, other dir=None)

Write the acqus dictionary in a file named 'filename.acqus'. Calls misc.write acqus 1D

#### **Parameters:**

• other\_dir: str or None
Different location for the acqus dictionary to write into. If None, self.datadir is used instead.

## write integrals(self, other dir=None)

Write the integrals in a file named '{self.filename}.int'.

#### Parameters:

• other\_dir: str or None

Different location for the integrals file to write into. If None, self.datadir is used instead.

## write procs(self, other dir=None)

Writes the actual procs dictionary in a file named 'filename.procs' in the same directory of the input file.

#### Parameters:

• other\_dir: str or None
Different location for the procs dictionary to write into. If None, self.datadir is used instead.
W! Do not put the trailing slash!

#### write ser(ser, acqus, path=None)

Writes a real/complex array in binary format. Calls misc.write\_ser. Be sure that acqus contains the BYTORDA and DTYPA keys. See misc.write\_ser to understand the meaning of these values.

#### **Parameters:**

- ser: *ndarray*Array that you want to convert in binary format.
- acqus: dict Dictionary of acquisition parameters. It must contain BYTORDA and DTYPA.
- path: strPath where to save the binary file.

#### xf1(self)

Process only the indirect dimension. Transposes the spectrum in hypermode or normally if FnMODE!= QF, then calls for processing.fp using self.procs[keys][0], finally transposes it back. The result is stored in self.S, then self.rr and self.ii are written. freq\_fl and ppm\_fl are assigned with the indexes of the transients.

## xf2(self)

Process only the direct dimension. Calls processing.fp using procs[keys][1] The result is stored in self.S, then self.rr and self.ii are written. freq\_f1 and ppm\_f1 are assigned with the indexes of the transients.