## week3

## August 23, 2023

Notebook file for Week 3, Diffusion experiment

Import libraries. The nmrbase folder needs to be located within the folder containing this Jupyter notebook

```
[]: import numpy as np
import scipy
import matplotlib.pyplot as plt
import nmrbase.expbase as expbase
import nmrbase.expdta as expdta
```

Diffusion NOTE: To analyze and report multiple datasets, simply copy the necessary blocks of code.

```
[]: filename = r"../DIRECTORY/FILENAME" #Defines the path to the data file. Here⊔

→ the path is relative to the current folder.
```

```
[]: a = expbase.expbase()
a.load(filename) # load data

f1=plt.figure()
ax1=f1.subplots()
a.plottm(ax1,1)
```

```
[]: ## TASKS:
## fine tune with set_xlim and set_ylim parameters to zoom in on the echo
ax1.set_ylim([VALUE, VALUE])
ax1.set_xlim([VALUE, VALUE])
f1
```

```
[]: a.pproc['digfmin']=1500  # set appropriate digital filter parameters
a.pproc['digfmax']=3500
a.digfilt()  # perform digital filter operation
```

```
[]: ## TASKS:
## fine tune with set_xlim and set_ylim parameters to zoom in on the echo
f2=plt.figure()
```

```
ax2=f2.subplots()
     a.plottm(ax2,1)
                         # plot the 1st scan
     ax2.set_ylim([VALUE, VALUE])
     ax2.set_xlim([VALUE, VALUE])
[ ]: ## TASKS:
     ## change parameter "ftmin" and "ftmax" to select the time interval \Box
      →corresponding to the echoes for the Fourier transform
     ## change parameter "ffmin" and "ffmax" to select the frequency range of the NMR_{\!lue}
      \hookrightarrow signal
     ## include statements to label axes
     ## use statements to change appearance such as font size, etc.
     f3=plt.figure()
     ax3 = f3.subplots()
     a.pproc['ftmin']=VALUE
                                                 # time interval for Fourier transform
     \hookrightarrow (0 = all data)
     a.pproc['ftmax']=VALUE
     a.pproc['ffmin']=VALUE
                                                  # frequency interval for spectrum_
      \rightarrow display (0 = all data)
     a.pproc['ffmax']=VALU
     a.pproc['dispper']=0.2
                                            # leave 20% space between each acquired_
     \hookrightarrow spectrum
     a.proc()
                                              # calculate Fourier transform of the data___
      \rightarrow in a
     ## this will use the digitally filtered data from before. Instead, the original \Box
      ⇒ data can be processed by loading it again.
     a.plotfrq(ax3,0)
                                              # plot the frequency domain data of the
      \hookrightarrow first scan from the data set
[]: # Gradient-duration-resolved integration of peaks on NMR spectra
     ## TASKS:
     ## change the intmin and intmax parameters to select the correct frequency range_{f U}
      \rightarrow for integration
     ## use set_xlabel and set_ylabel to set the labels to get a publication quality_
      \hookrightarrow figure
     f4=plt.figure()
     ax4=f4.subplots()
     a.pproc['intmin']=VALUE
                                     # set correct frequency range for integration
     a.pproc['intmax']=VALUE
```

Fitting of the integral to time axis. NOTE: For some results to fit, optimize the initial conditions

```
[]: # Fitting of the integral
     ## TASKS:
     ## change the initial fitting parameters to obtain proper fitting
     ## report the fitted gradient strength/diffusion coefficient with proper_
      ⇒significant figures
     ## use set_xlabel and set_ylabel to set the labels to get a publication quality_
     \hookrightarrow figure
     ## adjust appearance of figure as needed
     ax5 = plt.figure().subplots()
     a.idt.plot(ax5,disp=[0])
                                                # to be overlayed by fitted curve, __
     \rightarrow disp=0 plots only the NMR signal in trace 0
     big_delta=a.p.get('tau',0.1)
                                               # time interval between gradients
     def fun(t,a,b):
         return FORMULA
                                                  # INPUT the formula used for fitting.
     \rightarrow Use "np.exp()" for exponential, and t for small-delta (gradient duration).
     y = a.idt.dta[0]
     x = np.linspace(a.idt.x0,a.idt.x0+a.idt.dx*(a.pinc['n'][0]-1),a.pinc['n'][0])
     a.p1,a.p2=scipy.optimize.curve_fit(fun,x,y,p0=[A_VALUE,B_VALUE],maxfev=5000)
                     # SELECT the initial conditions, especially the exponential index
     print(round(a.p1[1],SF1),'±',round(np.linalg.eig(a.p2)[0][1]**0.5,SF2))
                   # REPLACE "SF1" and "SF2" with positive integrers to report the
      → fitting results with correct significant figures
```