F61/62: Nuclear Magnetic Resonance

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on Feburary 20th/ 21^{th} 2019

handed in as short report on: June 3, 2019

Abstract

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1 Introduction

1.1 Basics of Nuclear Magnetic Resonance

Nuclear Magnetic Resonance, or short NMR, can only be observed for molecules with a non-zero magnetic dipole moment. For a given Nuclei, whose spin \vec{J} is unequal to 0, one can calculate it's magnetic dipole moment as

 $\vec{\mu} = \hbar \gamma \vec{J}. \tag{1}$

Where γ is the gyromagnetic factor. In this experiment we will look at exclusively protons, which have $\gamma_{proton} = 2.6752 \cdot 10^8 sec^{-1} Tesla^{-1}$.

The magnetization of N nuclei can be obtained, by summing over all nuclei per unit volume

$$\vec{M} = \frac{1}{V} \sum_{i=1}^{N} \vec{\mu_i}$$
 (2)

In an external magnetic field the magnetic dipoles will realign themselves parallel or antiparallel to the direction of the magnetic field, cause of the interaction between \vec{B} and $\vec{\mu}$

$$\Delta E = -\vec{\mu} \cdot \vec{B_0} \tag{3}$$

With the assumption of a weak field $(\mu B \ll kT)$, one can approximate $\vec{M} \sim \frac{\vec{B_0}}{T}$ according to Marie Curie's Law.

In general, the magnetization can have an arbitrary direction relative to the external field. In the following we will decompose it into the components \vec{M}_{\parallel} (anti-)parallel and \vec{M}_{\perp} perpendicular to the external field. The magnetic dipole interacts with the \vec{B}_0 and as a result the general state of magnetization will dissipate it's excitation energy and reach the ground state asymptotically on a characteristic time scale.

This interaction results in a torque and since $\vec{B_0}$ is parallel to $\vec{M_{\parallel}}$, the torque only acts on $\vec{M_{\perp}}$.

$$\vec{\tau} = \vec{M} \times \vec{B_0} \tag{4}$$

Without a relaxation processes, the rate of change is given by

$$\frac{d\vec{M}_{\perp}}{dt} = -\gamma \vec{M}_{\perp} \times \vec{B}_{0}. \tag{5}$$

This differential equation can be solved by an Ansatz $\vec{M} = M_{\parallel}(\cos(\omega_L t), \sin(\omega_L t), 0)$ with ω_L being the Larmor frequency

$$\omega_L = \gamma B_0 \tag{6}$$

Let's consider the ground state magnetization \vec{M} , which is parallel to $\vec{B_0}$ and the z-axis. If we apply a sinusoidal voltage with frequency ω_{HF} to a coil which is coiled along the x-axis, it will result in a solenoidal magnetic field $\vec{B_1}$ which is perpendicular to $\vec{B_0}$. This will lead to \vec{M} precessing around $\vec{B_1}$. During a time interval Δt the angle α of the precession is then

$$\alpha = \gamma B_1 \Delta t. \tag{7}$$

If the time interval is chosen such that $\alpha = 90^{\circ}$, then \vec{M} is rotated into a perpendicular component $\vec{M_{\perp}}$ along the y-axis. Such a pulse is called a 90° pulse. Similarly we define 180° pulse which results in magnetization antiparallel to the static field $\vec{B_0}$.

1.2 Relaxation time

Now we want to consider the relaxation process, which can be described with the Bloch equations. Here we introduce the rotating frame of the transverse magnetization, where the transverse magnetization is constant, if no relaxation processes takes place. The Bloch equations assume that the time evolution is dominated by a restoring force which is proportional to the deflection from equilibrium

$$\frac{dM_{\perp}(t)}{dt} = -\frac{M_{\perp}(t)}{T_2} \tag{8}$$

$$\frac{dM_{\parallel}(t)}{dt} = -\frac{M_{\parallel}(t) - M_0}{T_1} \tag{9}$$

where T_2 is the spin-spin relaxation time, T_1 the spin-lattice relaxation time and M_0 the ground state magnetization.

In the laboratory system we can now write the equation (5) with (8) and (9) as

$$\frac{dM_{\perp}(t)}{dt} = -\frac{M_{\perp}(t)}{T_2} + \gamma(\vec{B} \wedge \vec{M})_{\perp} \tag{10}$$

$$\frac{dM_{\parallel}(t)}{dt} = -\frac{M_{\parallel}(t) - M_0}{T_1} + (\vec{B} \wedge \vec{M})_{\parallel} \tag{11}$$

1.2.1 Spin-spin relaxation time T_2

Spin-spin relaxation is caused by the interaction between the different magnetic dipoles and contributes to the transverse magnetization. This relaxation process is described with (10), which is be solved by

$$M_{\perp}(t) = M_{\perp}^{0} e^{-\frac{t}{T_{T_2}}}. (12)$$

There are 2 methods for measuring T_2 , which both use similar techniques. First let's look at the spin-echo method. We apply a 90° pulse at t=0 to generate a transverse magnetization \vec{M}_{\perp} . We then wait for a time $t=\tau$, after which we apply a 180° pulse and wait for till $t=2\tau$. At that time the magnetization should be transverse polarized again and we can measure the loss of amplitude. This described process is visualized in the following graphic.

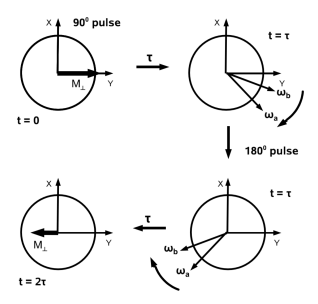


Figure 1: Dephasing of transverse magnetization $\vec{M_\perp}$

 ω_a and ω_b are denoting 2 different Protons which experience a different ω_L cause of the inhomogeneities in the external $\vec{B_0}$.

To measure the loss of amplitude we analyse the Fourier spectrum at the time $t=2\tau$ and integrate over the peak at the ω_L . If we repeat the process for different τ we will get a curve. This curve should follow (12), since the calculated amplitude is proportional to $\vec{M_{\perp}}$, and with the help of a fit we can extract the $T_{2,SE}$.

For the second method, the Carr-Purcell method, we also apply a 90° pulse at first and apply a 180° pulse at $t = \tau$. At $t = 2\tau$ the system should be in phase, but will dephase shortly after, same as in method one. We then apply another 180° at $t = 3\tau$ which will lead to a fully in phase system at $t = 4\tau$ again. This process can be continued for 180° at odd multiples of τ to get the system in phase at even τ . This process is a lot more precise for big τ in comparison to the spin-echo method, which gets inaccurate cause of the molecular diffusion (the protons move away from their original position) and inhomogeneities in the field. The measured amplitudes should still follow (12).

1.2.2 Spin-lattice relaxation time T_1

Spin-latice relaxation is caused by the interaction between the magnetic dipoles and the external magnetic field $\vec{B_0}$ and contributes to the parallel magnetization. This relaxation process is described with (11), which is solved by

$$M_{\parallel}(t) = M_0(1 - 2e^{-\frac{t}{T_{T_1}}}).$$
 (13)

We measure T_1 also with the spin-echo method. But this time we start at t = 0 with a 180° pulse and follow up after $t = \tau$ with a 90° pulse. We than measure the amplitude at $t = 2\tau$ and expect the resulting curve to follow (13).

1.3 Chemical shift

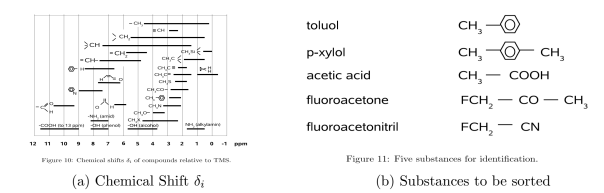


Figure 2: Chemical Shift

If a proton is bound by a molecule, it will experience a magnetic field that is weaker than the external magnetic field $\vec{B_0}$, since the electrons are shielding parts of it. This leads to a new resonance frequency ω_i for the proton bound by the molecule. This modified resonance frequency can be used to identify unknown substances. To do so, it is the easiest to compare it to a reference substance which you can easily identify in a Fourier Spectrum, since the whole frequency scale is dependant on $\vec{B_0}$ which means it's a relative scale and you can't work off absolute frequency values. We are using Tetra-Methyl-Silan or TMS since it will always be

clearly visible and in our case always be the peak on the most left. With the help of ω_{TMS} one can define a chemical shift

 $\delta_i = \frac{\omega_{TMS} - \omega_i}{\omega_L} \tag{14}$

for different molecule parts in reference to TMS as shown above in figure 2a. In this experiment we want to identify the substances in figure 2b from 5 different probes.

1.4 Imaging with NMR

Imaging with NMR is one of the most notable achievements of NMR. By adding a gradient to the external field $\vec{B_0}$ we create a coordinate dependency of the B_0 field, which implies a coordinate dependant ω_L . There are two different approaches to determine the density of the NMR active material of the probe in 1 dimension.

The first method is called frequency coding, where one has a set gradient over time and measures the the NMR signal in set times $t_n = n\Delta t$. Than one can do a discrete Fourier transformation of the N measurements and gets a finite amount of $M_{\perp}(n_i\Delta z)$ for the different coordinates.

The second method is called phase coding, where one has a set time and determines the phase of the precession all at the same time $t = t_0$. From that you again get a discrete amount of data points with which you can do a discrete Fourier transformation.

The main difference between these two methods is the time needed to record the data needed. The first method requires $T_f = N \cdot \Delta t$ compared to the $T_{\Phi} = t_0$ for the second method.

For 2 dimensions you have to use a combination of both since you need to determine 2 coordinates with just 1 signal. First a slice gets selected which is used as to derive the positional information. Then it does phase coding for a set gradient while measuring the NMR signal over time with a set time distance t_m . After finishing the frequency coding it changes the gradient, selects a slice again and does a phase coding again followed up by a frequency coding. It repeats this process now N different times for N different gradients over M different times. That leads to N × M data points which than can be transformed with a 2 dimensional Fourier transformation into the image matrix of the object. One of the N cycles is displayed in figure 3.

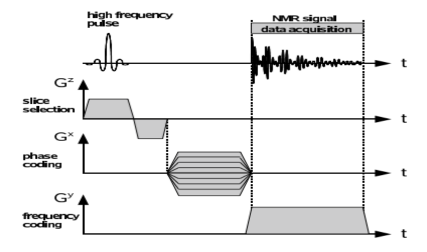


Figure 3: One of the N Loops needed for 2D Imaging

2 Layout of the experiment

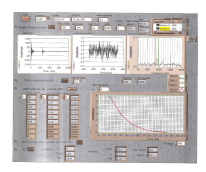
2.1 Relaxation Time and Chemical Shift

For the first two parts of the this experiment we use a minispec p20 which produces both of the magnetic fields we need for the relaxation time and measuring the chemical shift.

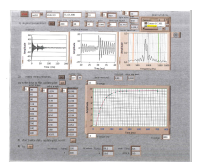


Figure 4: The minispec p20, electronic unit on the left, magnetic unit on the right

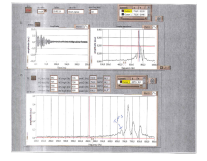
The constant $\vec{B_0}$, which is proportional to ω_L , can be adjusted by turning the screw sticking out of the Styrofoam as seen in Figure 4. The ω_{HF} will be set by the electronic unit of the p20 seen on the left side of Figure 4. An oscilloscope is also provided to measure the characteristic time for a 90° pulse, which is needed to calibrate the electronic unit of the p20 for the relaxation time measurements. The tubes with the substances get inserted into the magnetic unit parallel to the screw used to change ω_L . A high-pressure air nozzle can be used to rotate the tube to ensure a evenly distributed density of the substance in in the tube, we will use it for the chemical shift part since there only 1 measurement for each tube will be required.



(a) Ga $500 T_2$ Spin-Echo



(b) Ga $500 T_1$ Spin-Echo



(c) Chemical Shift

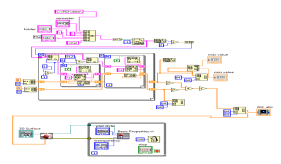
Figure 5: Example of the used Labviews

The data gets readout and analysed on a program written in Labview, the interface is shown in Figure 5 via some example measurements.

2.2 Imaging with NMR

We use a Bruker NMR analyzer 7.5 to generate the magnetic fields needed for 1D and 2D imaging. The readout gets fed into a computer where it gets analysed and can be later on displayed with a LabView .vi, examples of both can be seen in 6





(a) the Bruker NMR analyzer mq7.5

(b) LabView .vi to display 2d image

Figure 6: The NMR analyzer and LabView for 2D imaging

3 Measurements and evaluation

3.1 Measurement of relaxation time

We measure T_1 and T_2 once with the spin echo method and T_2 again with the Carr-Purcell sequence for each Ga 500 and Ga 600.

Substance	$T_{1,SE}$	$T_{2,CP}$	$T_{2,SE}$	
Ga 500	68.1	66.1	62.4	
Ga 600	99.2	84.0	78.6	

Table 1: Summary of relaxation time measurements in ms

We can see 3 relations from these measurements: $T_1 > T_2$ and $T_{2,CP} > T_{2,SE}$ for both substances as well as $T_{600} > T_{500}$ for each different method. The first can explained by The second relation can be explained by looking at the effects of the Carr-Purcell method. This method minimizes the effects of the molecular diffusion and field inhomogeneities, which improves the precision of greater echo times. The third relation can be attributed to the fact that the Ga 500 has higher concentration which leads to a more

From $\omega_L = 19.8 \text{ MHz} \cdot 2\pi$ we can also calculate our external field $\vec{B_0}$. And with the characteristic time $\Delta t = 1.29 \cdot 10^{-6}$ for a 90° pulse the solenoidal field $\vec{B_1}$ as well.

$$B_0 = \frac{\omega_L}{\gamma} = 0.48T \tag{15}$$

$$B_1 = \frac{\alpha}{\Delta t \gamma} = 4.3mT \tag{16}$$

where γ was the gyromagnetic factor mentioned at the beginning for protons.

3.2 Chemical shift

We measure the peaks and determine which peak is the TMS. After that we calculate the shift with the given ppm of the .vi and appoint them to a substance from figure 2b with the help of the provided cheat-sheet seen in figure 2a.

	A+	B+	C+	D+	E+
$\Delta(2\text{nd} - \text{TMS})$	2.2	2.1	2.0	3.9	2.6
$\Delta(3rd - TMS)$	3.9	6.9	11.6	6.3	7.5
$\Delta(4\text{th - TMS})$	6.3				
Substance	fluoroacetone	p-xylol	acetic acid	fluoroacetonitril	toluol

Table 2: Summery of chemical shift in ppm

Even though the resonance frequency of Flour is a lot higher than the frequency we are using here, we can see the peaks caused by the Flour. They appear because the of the spin-spin interaction between Flour and the proton(hydrogen) in FCH_2 which lead to 2 different states for the electrons. For both D+ and A+ we can see each of these peaks at 3.9 and 6.3 on both spectra.

From the width of these peaks we can additionally calculate the energy resolution of this measurement as well as the energy difference between the 2 different states FCH_2 can be in depending on the spin-spin interaction of the electrons.

$$\Delta E_{res} = f_{FWHM} \cdot h = 19.9Hz \cdot 4.136eVs = 8 \cdot 10^{-14}eV \tag{17}$$

$$\Delta E_{dipole} = f_{\Delta F} \cdot h = 48Hz \cdot 4.136eVs = 2 \cdot 10^{-13}eV$$
 (18)

3.3 Imaging with NMR

3.3.1 1d imaging

3.3.2 2d imaging

4 Discussion