500 MHz Solution-state NMR Procedure

(Bruker AVANCE Machines running TopSpin under WINDOWS XP)

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***Safety Issues***

⚠️ If you have metal implants, DO NOT do NMR yourself;

⚠️ Take everything ferromagnetic or vulnerable to magnetic field, such as mechanic watches, cellular phones, keys, credit cards, bank cards, tapes, computer disks, etc., out of your pockets and put them somewhere away from magnets;

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I. Sample preparation

1. NMR tubes of \textbf{5mm} in OD (typically 7” long) are used and available from:
   i. Chemistry Department Stockroom, UCSB (Phone: x2107)
   ii. Aldrich (Phone: 800-558-9160)
   iii. Wilmad (Phone: 800-220-5171).

   \textbf{Please get tubes for 500MHz or higher.}

2. Samples are dissolved in \textbf{deuterated} solvents for three purposes:

   Preferred concentration: >0.1 mM and >50mM for $^1$H and $^{13}$C, respectively, and sample volume: > 0.5 ml or >4 cm in height for 5 mm tubes.

   i. Deuteration removes solvent $^1$H signals which would otherwise dominate the $^1$H spectrum.
   ii. Deuterons provide a lock signal.

   Lock is a deuterium NMR process that the spectrometer uses to prevent the magnetic field from changing during the course of NMR experiments, thus locking the spectrometer.

   iii. Deuterons provide an internal reference for the spectra of $^1$H, $^{13}$C, $^{29}$Si, $^{31}$P, etc., rendering addition of reference standards such as TMS unnecessary.

   \textbf{Label your samples with your name and your advisor’s name. This helps us take care of unknown samples.}

II. Sign onto Logsheet

Enter

1. your name
2. your advisor’s name and department
3. your recharge account number (in the format: 8-4xxxxx-xxxxx-3)
4. your start time
5. \textbf{(Do this at the end of experiment:} your stop time and duration of experiment)
6. \textbf{(Do this at the end of experiment:} Status of instrument and report problems if any)

III. Start TopSpin Software

1. Make sure that the spectrometer is idle by looking at the computer. If yes, proceed to Step 2 below (if no, either wait, talk to the user on the machine, or do something else).

2. Login into the WINDOWS computer:
   Type your username, hit the Tab key (Don’t use return here!)
   Type your password, hit return
3. Double click the TopSpin icon on the desktop, the last dataset from your previous login session will appear.

   ![](icon.png)

   Power of right-click provides more functions and options. If you cannot find something you want, try the right mouse button.

**IV. How to Load/Change Samples**

1. Load samples onto the sample changer as below:
   i. Put samples in the blue spinners, measure depth with the depth gauge.
ii. Clean the bottom half of sample tube with napkin while holding the top half;

iii. Load them in the slots sequentially to the left of the white plastic piece with words “TO START:”, starting with the first slot.

iv. Please check that all samples have been correctly loaded and record the position of samples.

The rotation of sample tray is controlled by air either manually or automatically, depending on whichever mode is set and used. The rotation is clockwise when viewed from the top.
2. Push “Lift ON/OFF” button on the BSMS panel. The CDCl3 sample will come up and the sample tray will rotate. The 1st sample will be at the position to load and descend to the top of shim stack;

3. After ~10s (or when you don’t see the sample), press “Lift ON/OFF” button on the BSMS panel again to let the sample go down into the magnet. Wait for 10~15s for sample status LED to show DOWN (green);

4. Place the D2O sample in the next available slot following your sample(s), which would be the 1st slot if you have only one sample to run;

5. Run experiments using the procedure below for the 1st sample;

6. Once done with the 1st sample, repeat steps 2, 3, and 5 for the next sample and other samples if available;

7. Once done with all samples, do steps 2 and 3 to load the D2O sample, and lock the magnet for wrap-up as described in “Finishing up” at the end of the procedure.
V. $^1$H-NMR Setup and Data Acquisition

Click the flow chart below appears at the right side of the data area:

The guide will walk you through the data acquisition process interactively.

1. Click on New Experiment (or File $\rightarrow$ New) [New] (Words in brackets are the corresponding commands): a window pops up where you can
Name*: Soil Decomposition (e.g.) (meaningful or descriptive)  
EXPNO*: 1 (start with 1)  
PROCNO: 1 (start with 1)  
DU: C:\Bruker\TOPSPIN (don’t change)  
USER: nmrusu  
Solvent: CDC13 (leave it alone b/c it will be set when locking)  
Experiment: PROTON (experiment you want to do) or choose “Use Current Params” if you want to run the same experiment as the current data in display$^2$.  
Title: (any information useful for current sample, project, and/or experiment)  

*⚠️: IF YOU DO NOT CHANGE EITHER THE NAME OR THE EXPNO OF YOUR DATASET, YOU MAY OVERWRITE YOUR OLD DATA AND LOSE IT FOREVER.

$: To display an old dataset, go to the browser on the left, find and right-click on the desired dataset, and choose “Display” or “Display in a New Window”.

2. Click on Lock $^3$, which opens a solvents table. Select the solvent for your sample followed by OK. Wait for 1-2 min until you see that the Lock ON/OFF button on BSMS panel stays steady and a flat line (maybe noisy) sweeps back and forth in alternative red/green colors near the top of the lock window below. [See Appendix A: Lock troubleshooting].
3. Double click the small lock display screen to open the large lock display window. If it is not in the front, bring it up by clicking on the corresponding icon at the bottom of screen.
4. Check sample rotation (Don’t Click on **Sample Rotation** , just consider it a checklist item): **Make sure Sample Rotation is ON.**

   Press the **Spin ON/OFF** button on the BSMS panel to toggle sample rotation ON and OFF. When turning on, the button will be lit flashing first and then becomes steady in about 15 s.

5. **Shimming** (Don’t click on **Shim** , just consider it a checklist item):

   A) **Manual Shimming**

   - Make sure the lock display is visible. If not, bring it up by clicking on **Lock Display** at the bottom of screen or on the orange square at the right hand side of the upper toolbar.
   - Press “Z1” button and turn the wheel on the BSMS panel to shim Z1. The higher the lock signal, the better the shim. When the lock signal is at its highest, switch to Z2.
   - If lock signal is out of window, on the panel press the **Lock Gain** button and reduce the value by turning the wheel to bring lock signal down to the visible region of lock display.
   - Press “Z2” button and turn the wheel on the BSMS panel to shim Z2. The higher the lock signal, the better the shim. When the lock signal is at its highest, switch to Z.
   - Repeat Z1 and Z2 shimming above for 3 to 5 rounds until the lock level is optimized.
   - Press **Standby** button on the BSMS panel to deactivate the wheel, so as to prevent accidental change of shims.

B) Now you’re done with shimming and it is time to proceed for experiments.

   Please see **Appendix B: Manual Shimming** for how to shim manually.
6. Click on **Acquisition Pars** for modification of parameters. Set “TD” = 32k, “NS” = 8 and “DS” = 0 for simple $^1$H experiments for practical samples.

   Please refer to **Appendix** for simple spin dynamics of one-pulse FT-NMR.

![Image](image.png)

7. Click on **Prosol Pars.** to read the prosol (instrument dependent) parameters (Power levels, e.g. “PL1”, and Pulse lengths, e.g. “P1”).

8. Click on **Receiver Gain** and set Options to “Determine RG values automatically” [rga] based on the sample under study.

![Image](image.png)

After **OK** is clicked, wait for message “rga finished” in the status bar of TopSpin window to occur, indicating completion of rg setting.

9. Click on **Start Acquisition**. (If a warning message occurs, make sure to start acquisition in the right dataset). The FID display window below will appear along with some status parameters.
If parameters are not shown, right-click in the FID window, choose “Display Properties”, check “Status Parameters” and click on **OK**.

Three of the buttons near the top of the window may be used to:
- red square: Stop the acquisition [stop] with the data in buffer discarded
- black square: Halt the acquisition [halt] with the data in buffer saved to disk
- down triangle: Close the FID window

To monitor the acquisition status, just look at the Status Parameters to the right of FID. When Res. Time = 0, acquisition is completed and it is time to process the raw data. Or look at the bottom right of the Topspin status bar.

Click down triangle to close the FID window once acquisition finishes.
VI. Finishing Up

YOU ARE NOT FINISHED WITH THE SPECTROMETER UNTIL YOU DO THE FOLLOWING.

a. Make sure the D₂O sample is in the first slot of sample changer.
b. Press “Lift ON/OFF” button on BSMS panel to eject your sample and rotate the D₂O sample to the loading position.
c. In ~10s, hit “Lift ON/OFF” button on BSMS panel again to load the D₂O sample.
d. Wait for ~15s, click on \[\text{LOCK D₂O}\] to lock the spectrometer on the deuterated solvent and reset shim.

To lock, a dataset has to be displayed.
e. Quit lock window by pressing “Quit” or clicking X of the lock window.
f. Exit TopSpin by clicking the X at the top-right corner of the software.
g. Logoff your account: click Start (bottom-left corner) and choose logoff followed by “Yes”.
h. Important: On the logsheet, record your stop and duration times, and the spectrometer status. Report problems if any.
i. Important: remove all your samples from the tray and the NMR lab, and clean the lab space you have used.
VII. Data Processing Workstation

- Please go to NMR Processing Room (CNSI Room 1522) and use the computer denoted “Data Processing Workstation #2”

On the NMR Data Processing computers (WINDOWS XP, 128.111.243.67 or 68) (Feb. 2014)

1. Login
   a. Type “nmr_user” as a username and input password “4epp967”

2. Connect to NMR Data
   a. Click on “Start” at bottom left corner
   b. Choose “RUN”
   c. Type the desired machine name:
      i. Choose “\Nmr500sb-1” for solution state 500 MHz
      ii. Choose “\Varian-nmr” for solution state 600 MHz
      iii. Choose “\Nmr500wb-1” for solid state 500 MHz
      iv. Choose “\Ipso400wb” for solid state 400 MHz
      v. Choose “\Swbmri” for solid state 300 MHz
      vi. Choose “\Nmr800sb” for 800 MHz

3. Login using your own NMR account

4. A window will pop up displaying your NMR data folder (indicating a successful connection)

5. (OPTIONAL) Data Back-Up
   a. Go to folder: “Data”
   b. Find your personal directory and double click it
   c. Choose folder “nmr”
d. Now you can drag and drop the raw data files onto a flash/USB drive
6. Data Processing using TopSpin™
   a. Double click on “TopSpin 2.1” on desktop
   b. Find your desired machine name (e.g. “\nmr500sb-1” in the TopSpin™ browser panel on the left, expand it and navigate to your account folder
   c. (Optional) If you don’t see your desired machine name (e.g. “\nmr500sb-1” in the TopSpin™ browser panel on the left, add it yourself
      1. Right click in the browser panel area and choose “Add New Data Dir…”
      2. Type \nmr500sb-1 or \<IP address>
      3. Click “OK”
   d. Go to the dataset of interest by expanding \nmr500sb-1 and find your account name
   e. Right click on the dataset and choose “Display”
   f. Begin processing your NMR Data – please refer to the relevant NMR Procedure Manual
VIII. NMR Data Processing

*Click the TopSpin icon on the desktop to launch TopSpin
*Go to Menu Bar at the top and select: “Processing” → Choose: “Data Processing Guide”

![Diagram of NMR Data Processing Guide]

Please note that the first button Open Data Set can be skipped if you process the data just acquired and shown in the active window. Otherwise, use this button to open a different dataset for processing.

1) Click on Window Function and set “Line Broadening LB (Hz) =” 0.3 and “Window function type WDW =” exponential, followed by OK.
LB is a parameter to reduce noise level at the expense of resolution. The larger the LB, the better the S/N ratio but the worse the resolution. Typically “LB” = 0.3 Hz and the exponential WDW (Exp(-lb*t)) is multiplied with the raw data FID to generate a new FID with suppressed noise. [See Appendix D: LB and Window Function for details]

2) Click on Fourier Transform, and set “Standard Fourier Transform” “Size of real spectrum SI = 32768” and then click on OK to convert FID to spectrum (normally out of phase though).

To see the processed spectrum, click on Spectrum in the toolbar of data window.

3) Click on Phase Correction and check “Automatic Phasing” followed by OK [apk]. A phase corrected spectrum is obtained. (See Appendix C. Phase 1D Spectrum Manually for manual manipulation of phase correction)
4) **(Optional) Calibration:** this defines the position of 0 ppm, which is the chemical shift of TMS. Since the spectrum is referenced already to the solvent, chemical shift calibration is not necessary. However, if you do want exact calibration to TMS, click on button **Axis Calibration** and follow the on-screen instructions.

5) Click on **Baseline Corr.** and check “Auto-correct Baseline using Polynomial” [abs] followed by **OK** to correct the curvature and DC offset in the baseline of spectrum.

6) **(Optional) Advanced** : for multi-spectra display, add/subtract, and deconvolution.

7) Click on **Peak Picking**, choose “Define regions/peaks manually, adjust MI, MAXI”, and click on **OK**. In the data window, drag with the left mouse button a box (green) which defines MI (min. intensity), MAXI (max. intensity), by the bottom and top sides of the box, respectively, and chemical shift limits by the left and right sides. Peaks falling in this box both intensity- and shift-wise will be picked up and shown above the corresponding peaks.

![Peak Picking Image]

To modify the box, click on ![Modify Box] and drag sides or corners.  
To delete the box, click on ![Delete Box].  
To save the peak picking values, click on ![Save Values].

8) Click on **Integration** and choose “Define Integral Regions Manually” followed by **OK**. An integration window appears with the corresponding toolbar displayed at the top of spectrum:
If you see integration labels and values in the window opened up, delete them first by following the procedure “How to delete integral regions” below.

**How to Define Integral Regions and do Integration interactively:**

a. Click the following button (button turns green):
   Define integral region interactively

b. Put the red cursor line on one side of a peak or multiplet. (For accurate results, make sure the integration starts and ends at the baseline, and D1 is long enough).

c. Left-click-hold and drag the cursor line to the other side of the peak or multiplet.

d. Do step 2 and 3 for all regions to be defined and integrated.

e. Click the button to save integration and leave the integration mode.

**How to delete integral regions:**

- To delete all integral regions, click to Select/Deselect all integral regions, and click
- To delete a single region, right-click on the region to be deleted and choose “Select/Deselect” and then click on the delete button.

9) Click on Plot/Print and check “Print Active Window [prnt]”, or check “Print with Layout – start Plot Editor [plot]”, followed by OK. The former will print what you see in the TopSpin data display window while the latter uses the more sophisticated PlotEditor program for more controlled printing. (See Appendix E: Plot Editor for details.)
To Export Data, go to File → Export …, specify a folder in the “Look in” box, give a filename in the “File name” box with one of the “Legal File Extensions” shown in the “Files of type” box (e.g. 1H_5pEB.png), and click on the “Export” button.

10) (Optional) Click on E-mail/Archive to send the NMR data by email or archive it for storage. Since emailing data is prone to problems, archiving data is a better choice. In the E-mail/Archive window, choose “Save”.

Then select one of the choices in the next window (e.g. “Copy data set to a new destination” in this example or “Save data of currently displayed region in a text file”, which is a portable ASCII file) followed by OK.
Change DIR to the destination of your choice (e.g. E:, a flash drive) and click **OK**.

11) Wrap-Up
   a. Quit TopSpin™ software
   b. Logout of the “nmr_user” account

   **WARNING!!** If you don’t logoff, your NMR data could be at risk!
IX. Acquiring and Processing $^{13}$C NMR (1H decoupled)

Sometimes, an 1D $^{13}$C experiment may take hours or days depending on concentration (Recommended concentration is > 50 mM).

1. Go to Spectrometer $\rightarrow$ Data Acquisition Guide in the menu bar of TopSpin.
2. Click on New Experiment in the Guide window (see Procedure for $^1$H NMR).
3. In the New … window, choose one of the standard parameter files for $^{13}$C experiments below:

<table>
<thead>
<tr>
<th>Par file</th>
<th>Spectral information</th>
</tr>
</thead>
<tbody>
<tr>
<td>C13APT</td>
<td>CH+CH3 Positive, C+CH2 Negative (or vice versa), Intensity not quantitative</td>
</tr>
<tr>
<td>$^{13}$C PD</td>
<td>all carbons POSITIVE, for quantitative analysis</td>
</tr>
<tr>
<td>C13DEPT135</td>
<td>CH+CH3 Positive, CH2 Negative, C Gone, Intensity not quantitative</td>
</tr>
<tr>
<td>C13DEPT45</td>
<td>CH+CH2+CH3 Positive, C Gone, Intensity not quantitative</td>
</tr>
<tr>
<td>C13DEPT90</td>
<td>CH+CH3 Positive, C+CH2 Gone, Intensity not quantitative</td>
</tr>
</tbody>
</table>

The first two experiments are the most popular ones. If only the carbon types are of interest to you, use C13APT. If carbon signals are going to be quantified, choose C13CPD.

4. Click on Lock in the Guide if you have not done so (see Procedure for $^1$H NMR).
5. Click on Probe Match/Tune in the Guide, select “Automatic tuning / matching of ATM probe”, followed by OK. See Appendix I. Probe Tuning and Matching for Details.
6. Perform gradient shimming (see Procedure for $^1$H NMR).
7. Click on Acquisition Pars (see Procedure for $^1$H NMR) for modification of parameters. Set TD = 16k, NS = 64, DS = 0, and TD0 = 2000 for $^{13}$C experiments.
8. Click on Prosol Pars (see Procedure for $^1$H NMR).
9. Click on Receiver Gain (see Procedure for $^1$H NMR) and set Options to “Set RG values manually”. Input “16k” followed by return.

To know how long the experiment lasts beforehand, click on in the upper toolbar.
10. Click on Start Acquisition (see Procedure for $^1$H NMR).
11. Process data (see Procedure for $^1$H NMR).

A few things to note:

• You don’t have to wait until the experiment is done to process data. You can do processing as soon as the first NS scans are finished and saved to disk.

• A larger line broadening value is used for $^{13}$C than for $^1$H. Typically $lb = 1$ Hz.

12. Type "halt" if spectrum is satisfactory or let it continue if not. The acquisition will stop by itself after $NS*TD0$ number of scans are completed.

13. For data processing of a $^{13}$C spectrum, please refer to the procedure for $^1$H NMR.

14. Finish up as for $^1$H NMR.
X. Appendices

A. Lock troubleshooting

Problem: “Lock ON/OFF” won’t stop flashing and the lock signal stays at the bottom of the lock display window.

Possible reasons:
1. You started with a bad shimming file
2. Your sample is too concentrate, i.e. viscosity too high
3. Too little deuterons in your solvent

Solutions:
- for 1. Type [rsh] in the command line to read in a good shimming file and lock again
- for 2 & 3. Do manual locking as below:
  - Press “Lock ON/OFF” to turn off lock (LED off). At this point, you may see a noisy and weak oscillatory lock signal.
  - Hit “Lock Gain” and rotate the wheel on BSMS panel to increase lock signal amplitude until the signal fits the whole lock window.
  - Then press “Lock ON/OFF” to try to lock spectrometer. The LED should stay solid after brief flashing if spectrometer locks (the oscillatory signal becomes a flat sweeping line). If not,
  - Hit “Lock Power” and rotate the BSMS wheel to increase lock power. The lock signal will increase accordingly and may be saturated, i.e. the signal will not increase anymore with power. Remember: if the saturation happens, bring back the power by 2~3 units in reading.
  - Then press “Lock ON/OFF” and immediately increase “Lock Power” until the spectrometer is locked, where the “Lock ON/OFF” LED should stay solid and the oscillatory signal becomes a flat sweeping line. Avoid saturation stated above.
  - If you still have trouble, please ask Jerry/Jaya for help/debugging.

B. Manual Shimming (Don’t click on Shim , just consider it a checklist item).

- Make sure the lock display is visible. If not, bring it up by clicking on at the bottom of screen or on the orange square at the right hand side of the upper toolbar.
- Press “Z” button and turn the wheel on the BSMS panel to shim Z. The higher the lock signal, the better the shim. When the lock signal is at its highest, switch to Z2.
  - If lock signal is out of window, on the panel press the “Lock Gain” button and reduce the value by turning the wheel to bring lock signal down to the visible region of lock display.
Press “Z2” button and turn the wheel on the BSMS panel to shim Z2. The higher the lock signal, the better the shim. When the lock signal is at its highest, switch to Z.

- Repeat Z and Z2 shimming above for 3 to 5 rounds until the lock level is optimized.

- Press “Standby” button on the BSMS panel to deactivate the wheel, so as to prevent accidental change of shims.

### C. Phase 1D Spectrum Manually

Phase the spectrum (make all peaks absorptive)

1. Click on \( \mathcal{A} \) button (in the upper toolbar) and the toolbar above your spectrum should look like this:

   ![Toolbar](image)

2. Move the red cursor line to the tallest peak, right-click there, and select Set Pivot Point to mark the peak with a vertical red line, indicating a pivot point to be used by Ph1. Right-click again and choose “Calculate Ph0”. The tallest peak will now be phased roughly right.

3. Click and hold on \( 0 \) (zero order phase, used when the phase of peaks is frequency-independent). Dragging the mouse will change the zero order phase. Make the tallest peak exactly absorptive.

4. If other peaks are still not leveled on both sides, click and hold on \( 1 \) (first order phase, the phase of peaks is directly proportional to their frequency positions relative to the pivot point set above). Dragging the mouse to fix their phase. Many times this will also straighten out the baseline.

5. Click on \( \square \) to save the phase values and leave the phase mode.

6. If you want to process the same data again after you have done phasing, use “efp” (efp = em + ft + phase) instead of “ef”, so you don’t have to do “phase” again.
D. LB and Window Function

![Diagram showing LB and FT with different window functions and their corresponding FT results.]

E. Plot Editor

This is a very nice and user-friendly program. You are referred to the Plot Editor manual of TopSpin for more details and encouraged to use it often for more controlled printouts.

![Diagram of the Plot Editor interface with labels for selection modes, command buttons, major and minor editing modes, cursor, and status line.]
F. Spin Dynamics and Relaxation

Alignment of nuclei in magnetic field

Excitation, Acquisition and Relaxation

\[ 90^\circ \text{ RF} \quad \text{FID (Free Induced Decay)} \quad \text{Wait for } T_1 \]

\[ P1 \quad \text{AQ = TD*CW} \quad D1\sim3\sim5^* T_1 \]

\[ Z=Bo \quad \text{Mo} \quad y \]
G. Probe Tuning and Matching

When you do NMR on a nucleus X other than $^1$H and $^{13}$C (e.g. $X = ^2$H, $^{19}$F, $^{29}$Si, $^{31}$P, or ...), you will need to tune the probe. You do not normally need to do this when performing $^1$H or $^{13}$C NMR.

Steps

Click on **Probe Tune/Match** in the **Data Acquisition Guide** and select “Automatic tuning / matching of ATM probe”, followed by **OK**.

After ~45s, you will see a WOBB window showing the tuning/matching curve, the horizontal position of which corresponds to tuning and the depth to matching.

![WOBB Window](image)

The tuning curve should be aligned on the screen with the central redline and should reach all the way to the zero line of Y axis. If it doesn't look this way, then the probe isn't tuned.
The computer will tune and match the probe for you automatically. All you need to do at this point is to wait until you see the message “atma: finished” at the bottom left corner of Topspin.

H. Online NMR Book and Bruker NMR Encyclopedia

1) NMR Book: [http://www.cis.rit.edu/htbooks/nmr/](http://www.cis.rit.edu/htbooks/nmr/)
   Introduction to NMR concepts and practical issues.

   All you want to know about NMR.

I. Requirements for Access to the MRL NMR at CNSI

You have to pass the mini quiz within one month after training in order to be qualified for access to the NMR facility of MRL, which includes:

- Key Card for Lab & Building:
  1. Pass the MRL safety training;
  2. Fill out the CNSI access form:
     [http://www.cnsi.ucsb.edu/facilities/building_services/access/access_application.pdf](http://www.cnsi.ucsb.edu/facilities/building_services/access/access_application.pdf)
  3. Take the form to Sylvia in 2066G, MRL
- Web Scheduling Account (email Jerry for a setup appointment)
- NMR Account (email Jerry for a setup appointment)

These requirements apply to both on- and off-campus users.
XI. NMR Basic Principles

1. Spin

*Spin is a quantum mechanical phenomena that has no physical analog in classical physics. However, it will be helpful to visualize it as a small bar magnet that precesses about an axis.

*The existence of spin angular momentum is inferred by experiments, such as the Stern-Gerlach experiment, in which particles are observed to have angular momentum that cannot be solely accounted for by orbital angular momentum alone.

*Electrons, protons, and neutrons all have a value of spin +/- ½.
2. **Common NMR Nuclei**

<table>
<thead>
<tr>
<th>Nuclei</th>
<th>Unpaired Protons</th>
<th>Unpaired Neutrons</th>
<th>Net Spin</th>
<th>$\gamma$ (MHz/T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^1$H</td>
<td>1</td>
<td>0</td>
<td>1/2</td>
<td>42.58</td>
</tr>
<tr>
<td>$^2$H</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>6.54</td>
</tr>
<tr>
<td>$^{31}$P</td>
<td>1</td>
<td>0</td>
<td>1/2</td>
<td>17.25</td>
</tr>
<tr>
<td>$^{33}$Na</td>
<td>1</td>
<td>2</td>
<td>3/2</td>
<td>11.27</td>
</tr>
<tr>
<td>$^{14}$N</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>3.08</td>
</tr>
<tr>
<td>$^{13}$C</td>
<td>0</td>
<td>1</td>
<td>1/2</td>
<td>10.71</td>
</tr>
<tr>
<td>$^{19}$F</td>
<td>1</td>
<td>0</td>
<td>1/2</td>
<td>40.08</td>
</tr>
</tbody>
</table>

Larmor Frequency Equation:

$$\nu = \gamma B_o$$

where $\gamma$ is the gyromagnetic ratio (specific to each nucleus) and $B_o$ is the magnetic field strength.

3. **Energy Level Diagram**

![Energy Level Diagram](image)
4. cw NMR

5. Magnetization

*Alignment of nuclei in a magnetic field*
6. Pulsed NMR, Relaxation, and Detection

**Pulse Sequence**

- **D1 ~ 3*\(T_1\)**
- **P1**
- **FID (AQ = TD*DW)**

**Relaxation**
- \(T_1\) Relaxation
- \(T_2\) Relaxation

**Detector POV**
- 90° pulse