Multinukleární NMR spektroskopie C6800

- Jiří Pinkas, A12-224
- Materiály z přednášky v Isu
- Řešené úlohy ze spektroskopie nukleární magnetické resonance

http://nmr.sci.muni.cz

- Úlohy vyřešit a odevzdat
- Prezentace (na konci semestru) 10-15 min na vybrané téma NMR
- Závěrečná písemná zkouška

- 1922 Electron spin is observed (Stern-Gerlach)
- 1926 Nuclear spin David Dennison (H₂)
- 1938 I. I. Rabi observes NMR
 in a molecular beam of H₂
- Isidor I. Rabi awarded Nobel prize in physics 1944

"for his resonance method for recording the magnetic properties of atomic nuclei"





(1898 - 1988)

■ 1945 Purcell, Torrey, Pound @ Harvard

solid paraffin

■ 1945 Bloch, Hansen, Packard @ Stanford

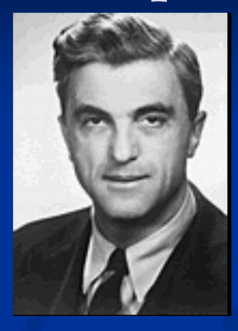
liquid H₂O

Varian Bros. & Russell

klystron for radars (WWII)

- 1948 Pake, van Vleck
- solid state NMR
- 1950 W. G. Proctor, F. C. Yu @ Stanford
 - δ chemical shift in ¹⁴NH₄¹⁴NO₃
- 1950 W. C. Dickinson @ MIT
- δ chemical shift in ¹⁹F
- 1952 Commercial NMR instruments used at DuPont, Shell, Humble Oil





Edward M. Purcell (1912-1997) & Felix Bloch (1905-1983)

NP in physics 1952

"for their development of new methods for nuclear magnetic precision measurements and discoveries in connection therewith"

- 1951 Proctor, Yu the first observed J scalar coupling ¹²¹Sb-¹⁹F in NaSbF₆
- 1951 Gutowsky, McCall, Slichter @ U. of IL J scalar coupling ³¹P-¹⁹F
- 1952 Hahn, Maxwell @ Berkeley J scalar coupling
- 1955 Bloom, Shoolery spin decoupling
- 1960 Shoolery integration
- 1966 Ernst, Anderson FT NMR at Varian
- 1968 Waugh @ MIT HR, multipulse NMR in solids

■ 1971 Jeener - 2D NMR

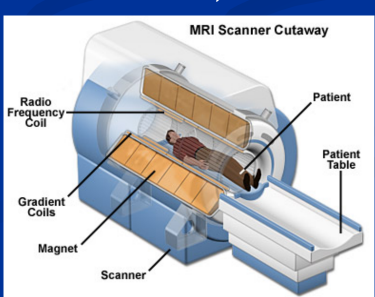
1971 Damadian - different NMR relaxation times of tissues and tumors

■ 1972 CP, HP decoupling

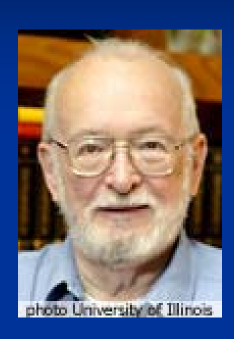
■ 1972 The first routine ¹³C NMR spectrometer

(before mainly ¹H, ¹⁹F, and ³¹P NMR)

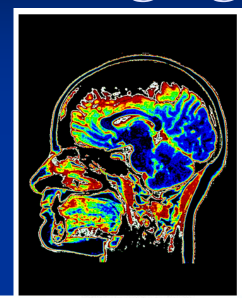
1973 Lauterbur - MRI

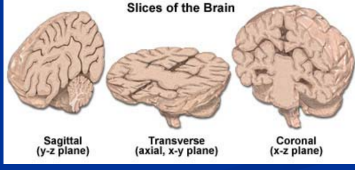


MRI-Magnetic Resonance Imaging



Paul C. Lauterbur (1929-)







Sir Peter Mansfield (1933-)



NP in physiology and medicine 2003

■ 1974/1979 R. R. Ernst 2D COSY, NOESY

■ 1977 MAS

■ 1981 Bax, Freeman INADEQUATE

■ 1982 APT

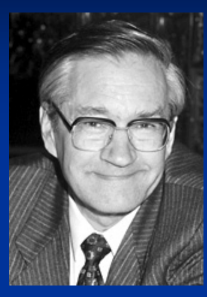
■ 1983 Freeman BB decoupling, MLEV, WALTZ

■ 1990 3D and ¹H/¹⁵N/¹³C Triple resonance

■ 1991 R. R. Ernst NP in chemistry

2001 The first commercial 900 MHz instrument

2002 K. Wüthrich NP in chemistry

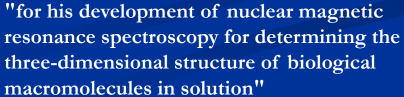


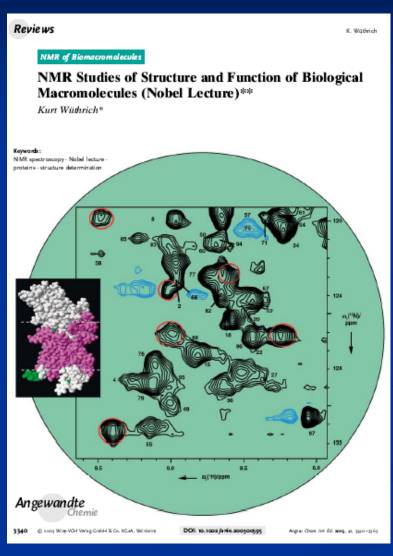
Richrad R. Ernst (1933-) NP in chemistry 1991

"for his contributions to the development of the methodology of high resolution nuclear magnetic resonance (NMR) spectroscopy"



Kurt Wüthrich (1938-) NP in chemistry 2002



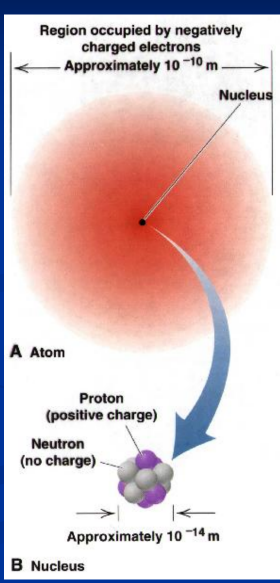


Nuclear Magnetic Resonance

- High resolution liquid state NMR spectroscopy
- Solid state NMR spectroscopy
- High-pressure NMR
- NMR in the gas phase
- NMR spectroscopy in liquid crystalline media
- Magnetic resonance imaging (MRI)

Hyperfine Interactions

- Interactions of nuclei with the electric and magnetic fields
- Interactions between a nucleus and electrons
- Transfer of chemical (electronic) information from bonds and lone pairs to a nucleus:
 - Indirect
 - Direct



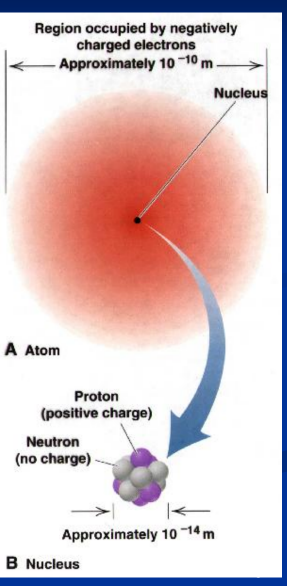
Hyperfine Interactions

Indirect

- Electric field gradient (EFG) with nuclear electric quadrupole
- Induced magnetic field with nuclear magnetic moments (shielding)

Direct

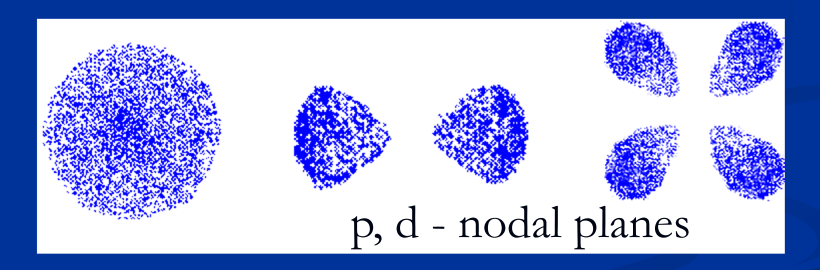
• s-electrons within nuclei, polarization of bonding spins by nuclear spin (J-coupling)



Direct Interactions

ONLY s-electrons can interact with nuclei

ONLY s-electrons have non-zero electron density at a nucleus



Which quantum number determines the number of angular nodes? Which quantum number determines the number of radial nodes?

Relationship Between Wavelength, Frequency and Energy

- Speed of light (c) is the same for all wavelengths $c = 2.9979 \ 10^8 \ \text{m s}^{-1}$
- Frequency (ν), the number of wavelengths per second, is inversely proportional to wavelength:

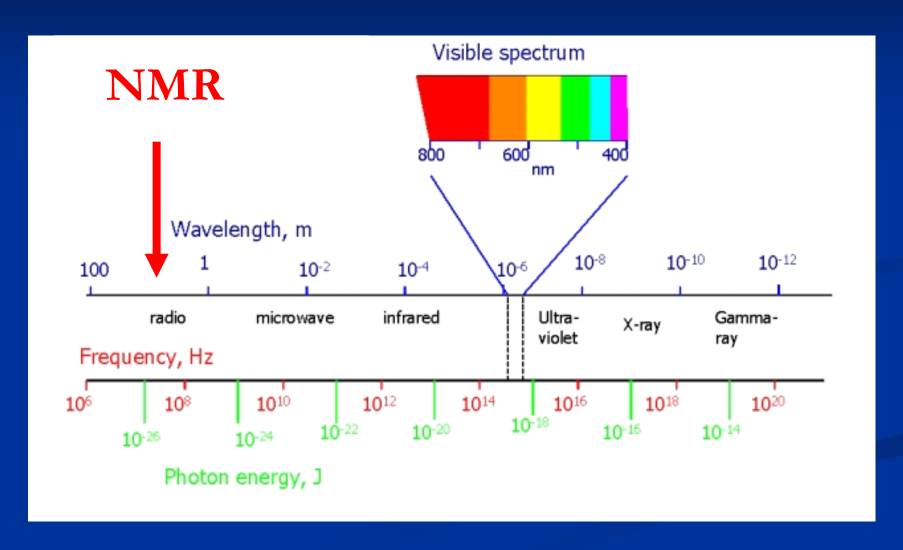
$$v = c/\lambda$$

Energy of a photon is directly proportional to frequency and inversely proportional to wavelength:

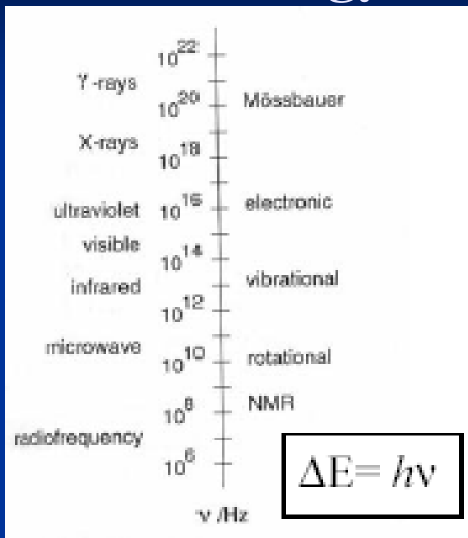
$$E = hv = hc/\lambda$$

h = Plank's constant = 6.626176 10⁻³⁴ J s

Electromagnetic Radiation



Method Energy Scale



Energy Scale Conversion Factors

	Hz	eV	J mol ⁻¹
Hz	1	4.136 10 ⁻¹⁵	3.990 10 ⁻¹⁰
eV	2.418 10 ¹⁴	1	9.649 10 ⁴
J mol ⁻¹	2.506 10 ⁹	1.036 10 ⁻⁵	1

Isotopes

Isotopes = a set of nuclides of an element, same Z, different A there is about 2600 nuclides (stabile and radioactive)
340 nuclides found in nature
270 stabile and 70 radioactive

Monoisotopic elements: ⁹Be, ¹⁹F, ²³Na, ²⁷Al, ³¹P, ⁵⁹Co, ¹²⁷I, ¹⁹⁷Au

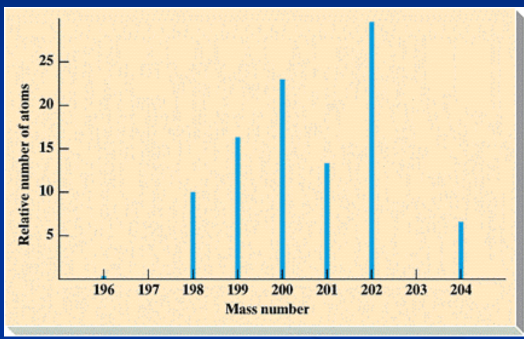
Polyisotopic elements: ¹H, ²H (D), ³H (T) ¹⁰B, ¹¹B Sn has the highest number of stabile isotopes – 10

112, 114, 115, 116, 117, 118, 119, 120, 122, 124Sn

Natural Abundance, %

Isotopic Compositions of the Elements

^A Hg	I	NA%
196	0	0.146
198	0	10.02
199	1/2	16.84
200	0	23.13
201	3/2	13.22
202	0	29.80
204	0	6.850



Mass number, A

Natural Abundance, %

Isotopic Compositions of the Elements

¹H	99.985	¹⁶ O	99.759
^{2}H	0.015	¹⁷ O	0.037
		¹⁸ O	0.204
¹² C	98.89		
¹³ C	1.11	³² S	95.00
		33 S	0.76
¹⁴ N	99.63	³⁴ S	4.22
¹⁵ N	0.37	³⁶ S	0.014

Variability in Isotopic Compositions

Natural Abundance, %

Isotope	Range	Average
$^{10}\mathrm{B}$	18.927 - 20.337	19.9 (7)
$^{11}\mathbf{B}$	81.073 - 79.663	80.1 (7)
¹⁶ O	99.7384 - 99.7756	99.757 (16)
¹⁷ O	0.0399 - 0.0367	0.038 (1)
¹⁸ O	0.2217 - 0.1877	0.205 (14)

```
electron spin s = \frac{1}{2}
```

proton and neutron
$$I = \frac{1}{2}$$

nuclear spin
$$I = z^{1/2}$$
 $z = integer 0, 1, 2, 3,$

Number of protons, Z	Number of neutrons, N	I
even	even	0
odd	odd	integer
even	odd	
odd	even	multiples of $\frac{1}{2}$

protons and neutrons are Fermions, obey Pauli exclusion principle

¹²C

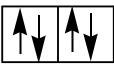


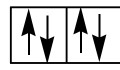


13**C**

 $I = \frac{1}{2}$

$$I = 0$$





$$\uparrow$$

$$\downarrow$$

$$\mathbf{n}$$

p

- even even: I = 0 ⁴He, ¹²C, ¹⁶O, ²⁰Ne, ²⁴Mg, ²⁸Si, ³²S, ³⁶Ar, ⁴⁰Ca
- odd odd: *I* = integer
 ONLY ²H, ⁶Li, ¹⁰B, ¹⁴N, ⁴⁰K, ⁵⁰V, ¹³⁸La, ¹⁷⁶Lu
- even odd and odd even:

 $I = \text{multiples of } \frac{1}{2}$ ¹³C ¹/₂, ¹⁷O 5/2, ³³S 3/2

Number of protons Z	Number of neutrons N	Number of nuclides 168 8 50		
even	even	168		
odd	odd	8		
odd	even	50		
even	odd	57		

Allowed Nuclear Multipole Moments as a function of Spin I

	l = 0	l = 1	l=2	l = 3	l=4
Spin	monopole	dipole	quadrupole	octapole	hexadecapole
I=0	electric	0	0	0	0
$I = \frac{1}{2}$	electric	magnetic	0	0	0
I = 1	electric	magnetic	electric	0	0
$I = \frac{3}{2}$	electric	magnetic	electric	magnetic	0
I=2	electric	magnetic	electric	magnetic	electric

Nuclear Magnetic Dipole Moment couples to magnetic field

Ν



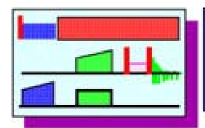
5

H	ļ ;			pin	_												Не
Li	Be		S	pin	> 1/2							В	C	N	O	F	Ne
Na	Mg											Al	Si	P	S	Cl	Ar
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
Rb	Sr	Y	Zr	Nb	Mo	Tc	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	I	Xe
Cs	Ba	La	Hf	Ta	\mathbf{w}	Re	Os	Ir	Pt	Au	Hg	Tl	Pb	Bi	Po	At	Rn
Fr	Fr Rd Ac Ce Pr Nd Pm Sm Eu Gd Th Dy Ho Er Tm Yh Lu																

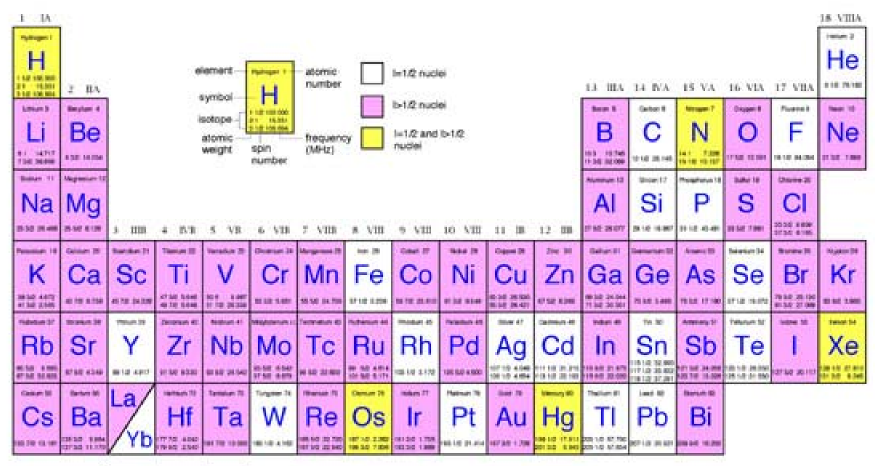
Th Pa U Np Pu Am Cm Bk Cf Es Fm Md No Lr

Nuclear Electric Quadrupole Moment couples to electric field gradient





Elements Accessible by NMR





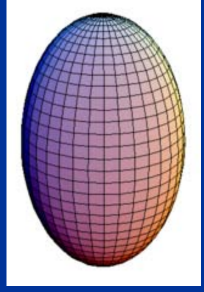
NO stable nucleus has spin 2

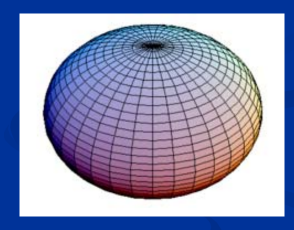
the highest value of spin for a stable nucleus is 7

unstable nuclei
highest integral spin 16 - isomer ¹⁷⁸Hf
highest half-integer 37/2 - isomer ¹⁷⁷Hf)

- Nuclei with spin ½ a spherical charge distribution
- Nuclei with $I > \frac{1}{2}$ nonspherical charge distributions

(prolate or oblate)





- Nuclei with a non-zero spin \rightarrow magnetic moment (μ)
- Nonspherical nuclei \rightarrow electric quadrupole moment (e Q)

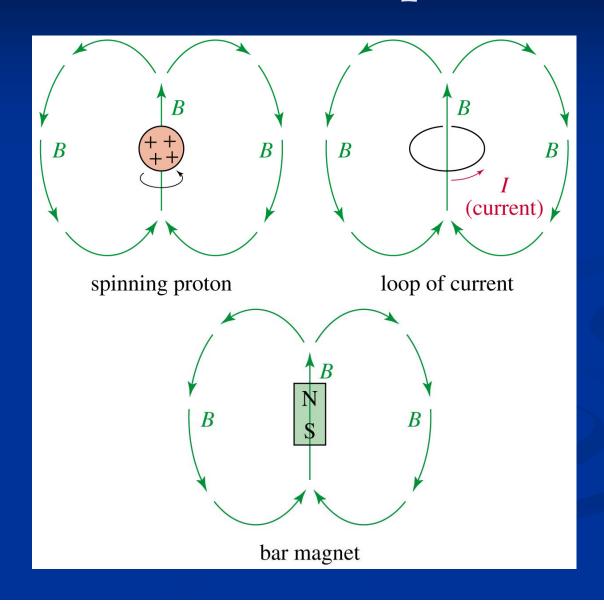
Properties of Nuclei:

Mass, Charge, Spin and Magnetism.



"The concept of spin is difficult. It was forced upon scientists by the experimental evidence". Malcolm Levitt.

Rotating positive charge generates magnetic field



Nuclear spin = Spin angular momentum, P (vector)

(moment hybnosti)

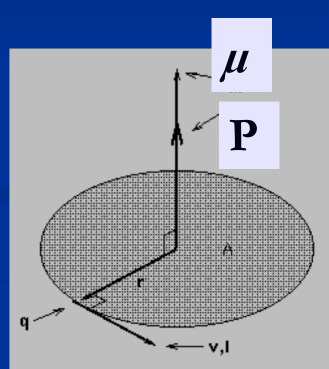
Spin quantum number IMagnetic quantum number $m_{\rm I}$

Magnitude of *P* is quantized:

$$P = \frac{h}{2\pi} \sqrt{I(I+1)}$$

Direction with respect to the magnetic field B_0 is quantized:

$$P_z = \frac{h}{2\pi} m_I$$



Spin Angular Momentum, P

$$P = \frac{h}{2\pi} \sqrt{I(I+1)} \qquad P_z = \frac{h}{2\pi} m_I$$

$$P_z = \frac{h}{2\pi} m_I$$

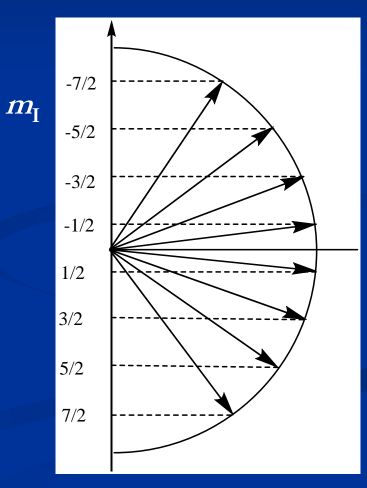
I = Nuclear spin quantum number $I = 0, \frac{1}{2}, 1, \frac{3}{2}, \frac{5}{2}, \frac{3}{7}, \frac{7}{2}, \dots$

 $m_{\rm I}$ = Nuclear spin magnetic quantum number

Multiplicity, M = 2I + 1 values $m_{I} = I, I - 1, I - 2, ..., -I + 2, -I + 1, -I$

$$\cos \theta = \frac{P_z}{P} = \frac{m_I}{\sqrt{I(I+1)}}$$

 59 Co, I = 7/2



Spin Angular Momentum, P

$$P = \frac{h}{2\pi} \sqrt{I(I+1)}$$

I	$[I(I+1)]^{1/2}$
1/2	0.866
1	1.414
3/2	1.936
5/2	2.958
3	3.464
7/2	3.969
4	4.472
9/2	4.975

Spin Magnetic Moment, µ

The electrons, nucleons (protons, neutrons) and some nuclei possess intrinsic magnetism, which is not due to a circulating current.

Permanent magnetic moment similarly as spin angular momentum.

Magnetic moment, μ , is directly proportional to the spin angular momentum, P:

$$\mu = \gamma P$$

γ is the gyromagnetic (magnetogyric) ratio

Magnetogyric Ratio

 γ - the magnetogyric ratio is the ratio of the nuclear magnetic moment μ to the nuclear angular momentum **P**.

$$\mu = \gamma P$$

γ - Important characteristic of nuclei

$$[rad T^{-1} s^{-1}]$$

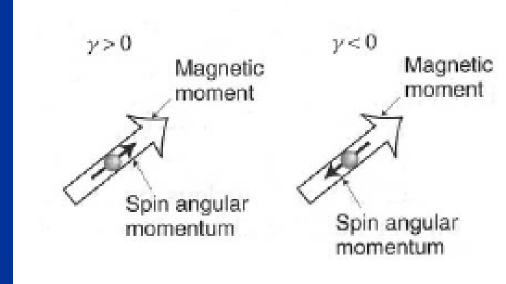
Spin Magnetic Moment, μ $\mu = \gamma P = \gamma \hbar [I(I+1)]^{1/2}$

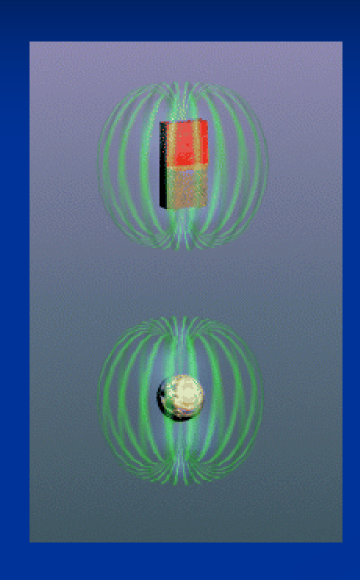
$$\mu_{\rm z} = \gamma P_{\rm z} = \gamma \hbar m_{\rm I}$$

 ^{2}H 13**C** 15N 19**F** ²⁹Si 31**p** Nucleus ^{1}H $\gamma [10^{-7} \text{ rad } T^{-1} \text{s}^{-1}]$ 26.75 4.11 6.73 -2.7125.18 -5.3210.84

electron

$$\gamma_e = 17 609 10^7 = 658 \gamma(H)$$

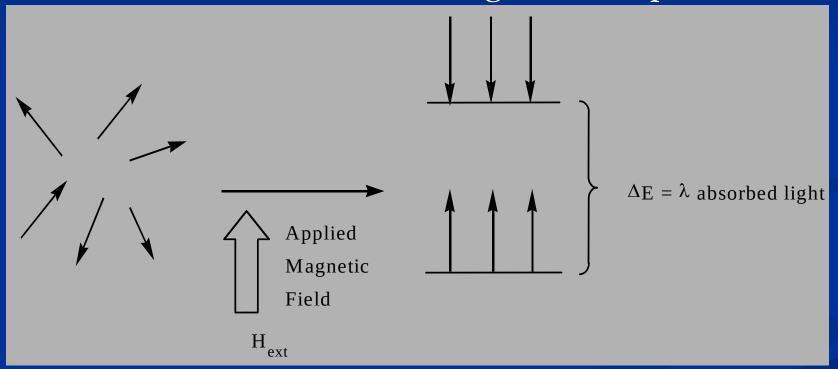






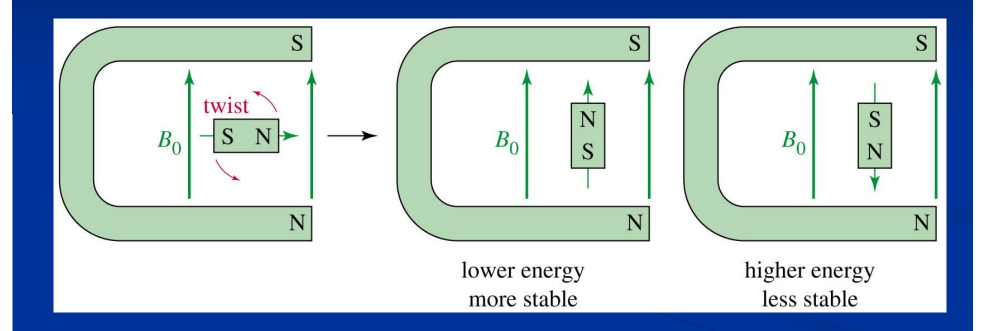
Random orientation

Zeeman plitting to 2I + 1 levels Alignment of spins



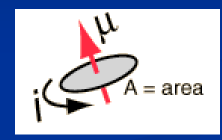
No Field

Magnetic Field



magnetic dipole

- An angular momentum is associated with each rotating object
- A nuclear spin possesses a magnetic moment μ arising from the angular momentum of the nucleus



• The magnetic moment μ is a vector perpendicular to the current loop

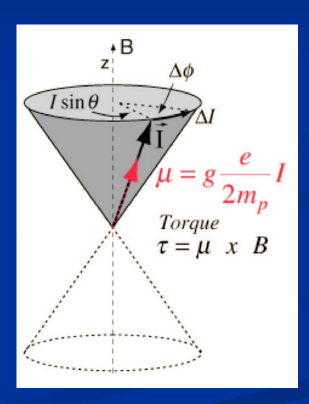
$$\mu = iA$$

• In a magnetic field (B) the magnetic moment behaves as a magnetic dipole

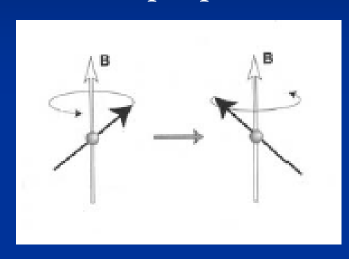
In B_0 , a magnetic moment μ is directed at some angle w.rt. B_0 direction

the B_0 field will exert a torque on the magnetic moment. This causes μ to precess about the magnetic field direction

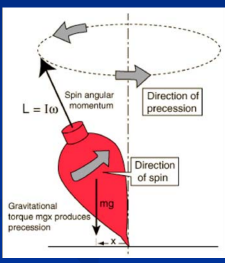
Torque is the rate of change of the nuclear spin angular momentum



Spin precession in the external magnetic field.







Quantum description of precession shows that the frequency of the motion is:

$$\omega_0 = -\gamma B_0 \text{ [rad s}^{-1} \text{] or } v_0 = -\gamma B_0 / 2\pi \text{ [Hz]}$$

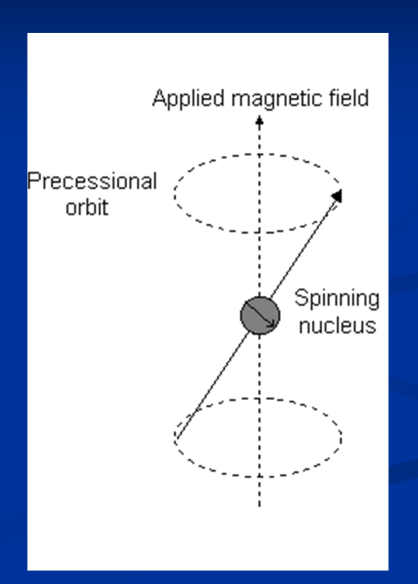
It is called the Larmor frequency (if $\gamma > 0$ then $\nu_0 < 0$)

Larmor Frequency

$$\omega_0 = -\gamma B_0 \text{ [rad s}^{-1}]$$

$$v_0 = -\gamma B_0 / 2\pi [Hz]$$

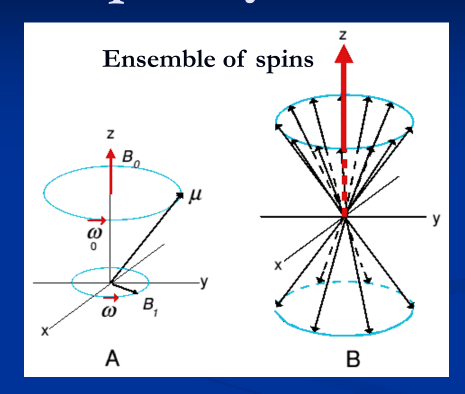
$$\nu_0 = -\frac{\gamma B_0}{2\pi}$$



Larmor Frequency

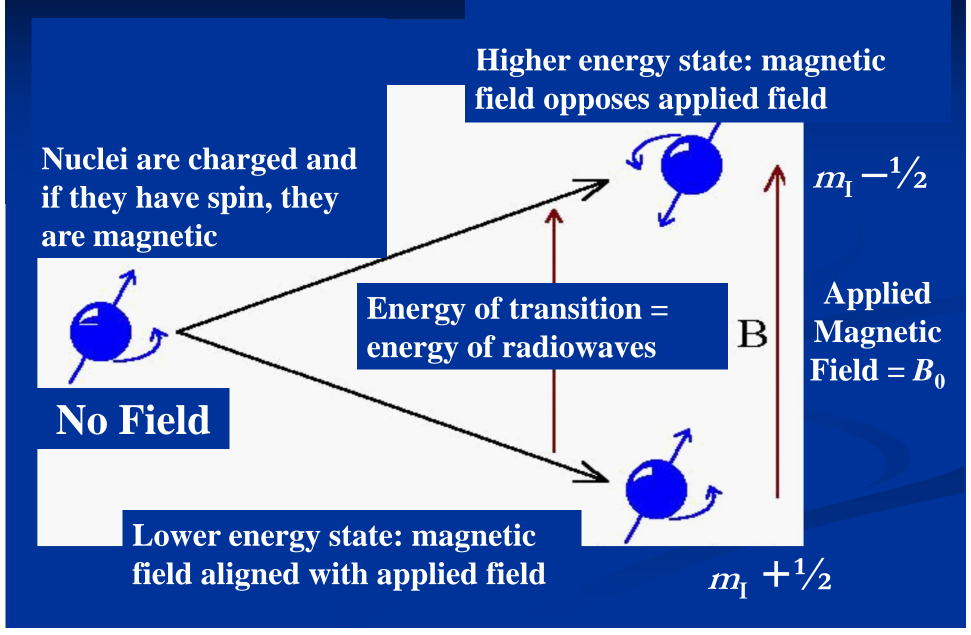


Sir Joseph Larmor (1857-1942)



$$\omega_0 = -\gamma B_0 \text{ [rad s}^{-1} \text{] or } v_0 = -\gamma B_0 / 2\pi \text{ [Hz]}$$

Nuclear Zeeman Effect - Splitting



The magnetic energy depends on the interaction between the magnetic moment and B_0 field:

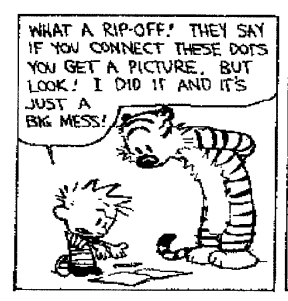
$$E_{\text{mag}} = -\mu \cdot B_0$$
 (a scalar product of 2 vectors)

$$E_{\text{mag}} = -\mu_{z} B = -\gamma P_{z} B$$

$$E_{\text{mag}} = -m_{\text{I}} \hbar \gamma B$$

NMR selection rule $\Delta m_{\rm I} = \pm 1$

CALVIN AND HOBBES By Bill Watterson





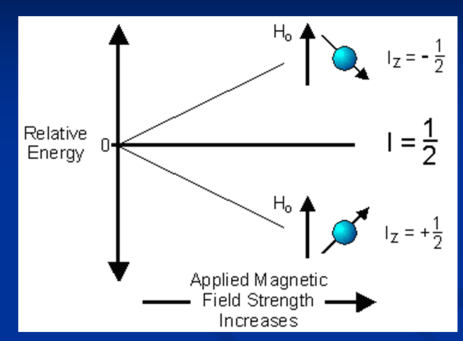




Spin in Magnetic Field

$$I = \frac{1}{2}$$
 $E_{\text{m}=-1/2}$

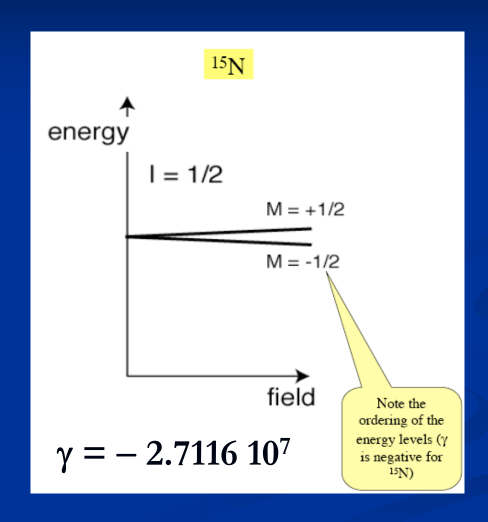
 $|E|_{\mathrm{m}=1/2}$



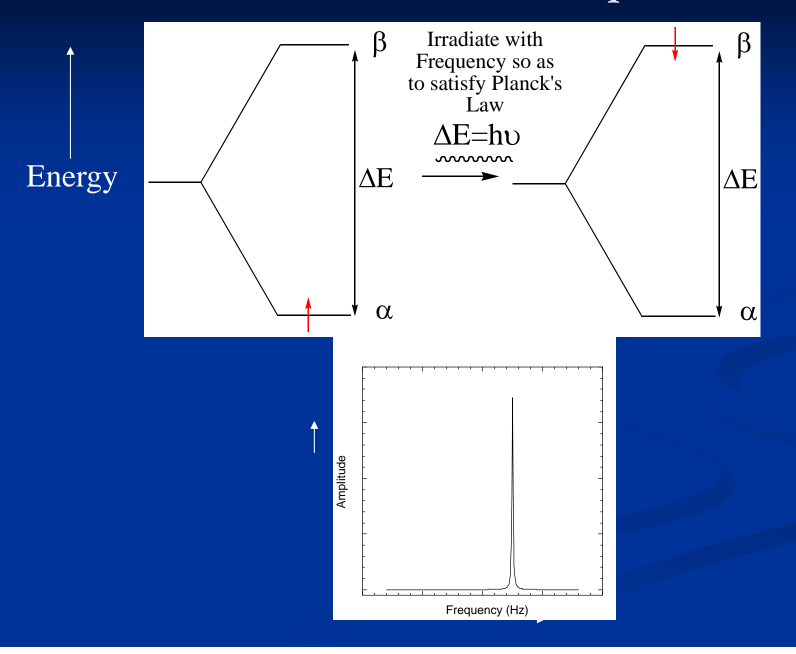
$$\Delta E_{\text{mag}} = E_{\text{m=-1/2}} - E_{\text{m=1/2}} = \Delta m_{\text{I}} \hbar \gamma B = \hbar \nu \Rightarrow \nu = \gamma B/2\pi$$

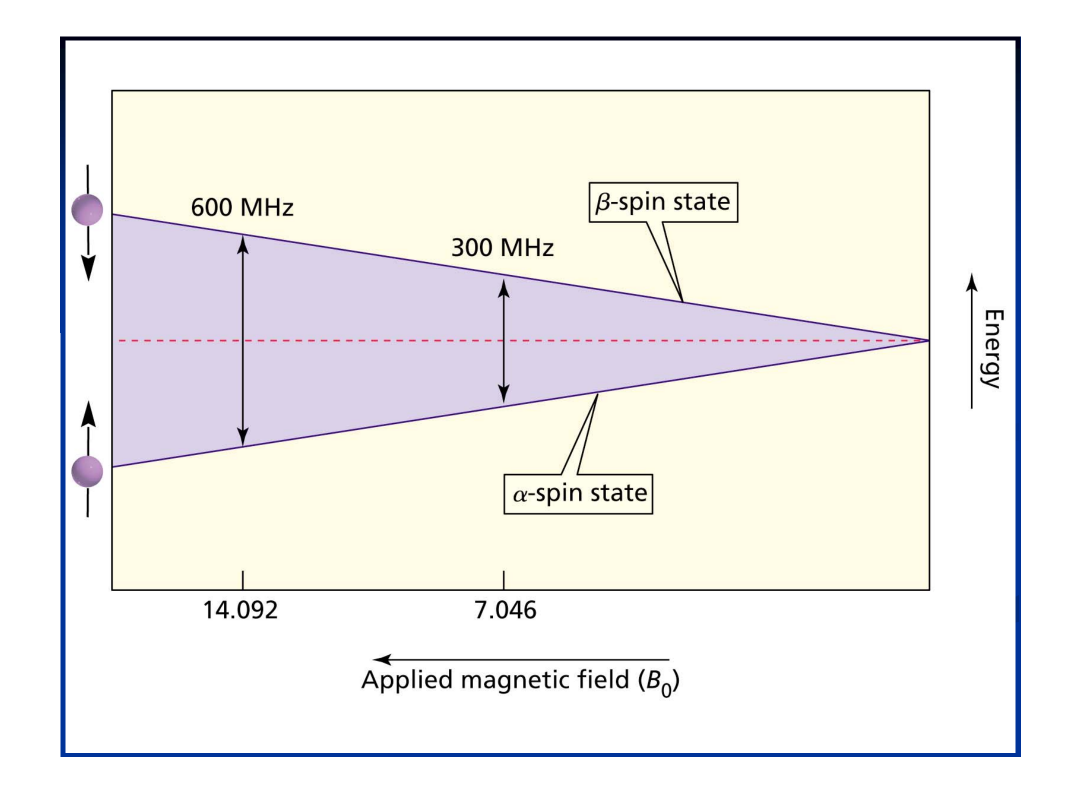
The frequency of the electromagnetic radiation that corresponds to the energy difference between the two energy levels is equal to the precessional frequency of the nuclei.

Spin in Magnetic Field



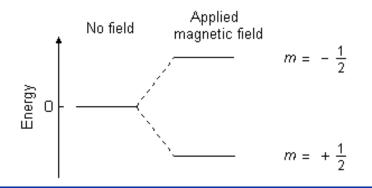
Excitation of NMR Spin

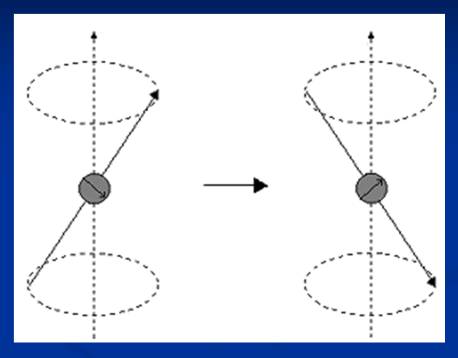




Energy Levels for $I = \frac{1}{2}$

Energy levels for a nucleus with spin quantum number 1/2



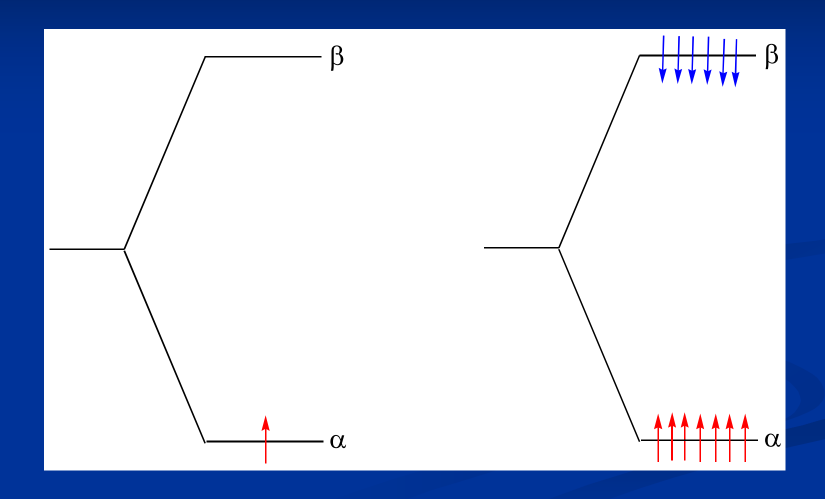


$$\Delta E_{\text{mag}} = E_{\text{m=-1/2}} - E_{\text{m=1/2}} = \Delta m_{\text{I}} \hbar \gamma B = \hbar \gamma B/2\pi$$

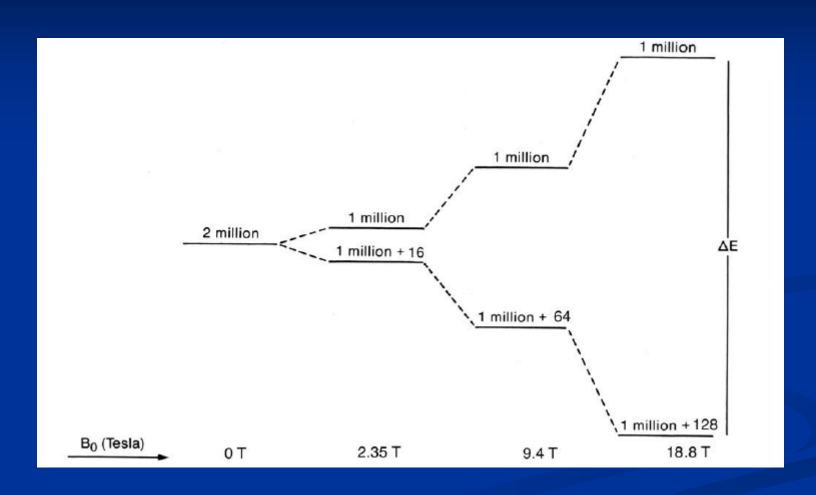
Protons

 $\Delta E = (6.626 \ 10^{-34} \ J \ s \ 26.75 \ 10^7 \ rad \ T^{-1} s^{-1} \ 11.743 \ T)/2\pi = 3.313 \ 10^{-25} \ J$ very small energy difference

Energy Levels for $I = \frac{1}{2}$



Energy Levels for $I = \frac{1}{2}$



Boltzmann Distribution

The excess of nuclei on the lower energy level is given by Boltzmann distribution:

$$\frac{N \uparrow \downarrow}{N \uparrow \uparrow} = \exp\left(-\frac{\Delta E}{k_B T}\right) = \exp\left(-\frac{\hbar \gamma B}{k_B T}\right) =$$

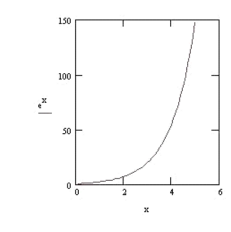
 $= \exp(-3.313 \ 10^{-25} / \ 4.101 \ 10^{-21}) = \exp(-8.078 \ 10^{-5}) = 0.99991922$

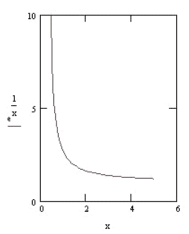
If $N\uparrow\uparrow = 1\ 000\ 000$ then $N\uparrow\downarrow = 999919$ Only 81 out of 2 million ¹H nuclei contribute to NMR signal at 500 MHz!

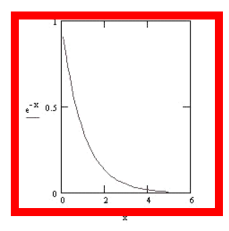
 $h = 1.055 \ 10^{-34} \ J \ s$ $\gamma_H = 26.75 \ 10^7 \ rad \ T^{-1}s^{-1}$ $B = 11.7433 \ T \ (500 \ MHz)$ $k_B = 1.3807 \ 10^{-23} \ J \ K^{-1}$ $T = 297 \ K$

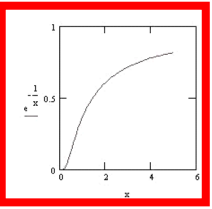
Boltzmann Distribution

 $N\uparrow\downarrow/N\uparrow\uparrow = \exp(-\Delta E/k_B T) = \exp(-\hbar \gamma B/k_B T)$









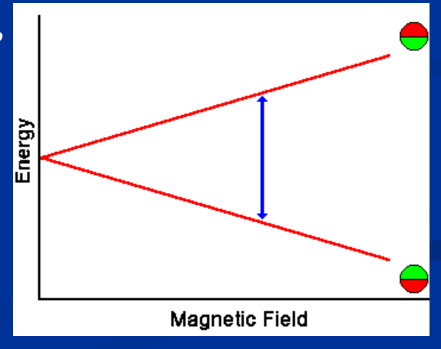
the stronger the field and the higher the magnetogyric ratio, the larger the population difference

the higher the temperature, the smaller the population difference

Boltzmann Distribution

The higher the field B, the larger the energy difference, the larger the population difference,

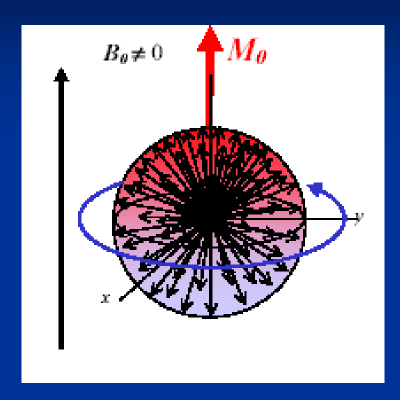
the larger the net magnetization, and the bigger the NMR signal



Nuclear Magnetic Resonance (NMR)

- Nuclear spin ½ nuclei (e.g. protons) behave as tiny bar magnets.
- Magnetic a strong magnetic field causes a small energy difference between $+ \frac{1}{2}$ and $-\frac{1}{2}$ spin states.
- Resonance photons of radio waves can match the exact energy difference between the $+\frac{1}{2}$ and $-\frac{1}{2}$ spin states resulting in absorption of photons as the protons change spin states.

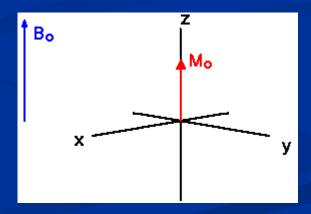
Magnetization



More nuclei point in parallel to the static magnetic field.

The macroscopic magnetic moment, M_0

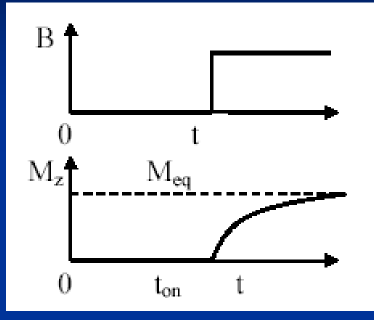
$$M_0 = \sum \mu_i$$



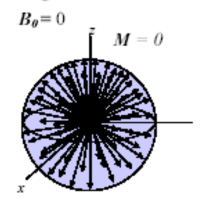
In-Field

Bloch equations: the nuclear magnetization $M = (M_x, M_y, M_z)$ as a function of time and relaxation times T_1 and T_2

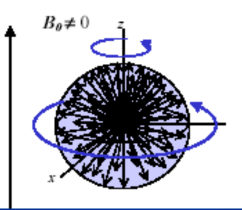
Longitudinal Magnetization



No magnetic field

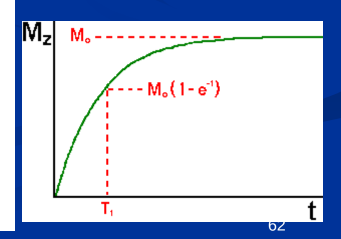


Magnetic field



$$\frac{d}{dt}M_{Z} = R_{1}(M_{eq} - M_{z}) = \frac{1}{T_{1}}(M_{eq} - M_{z})$$

$$M_z(t) = M_{eq}(1 - e^{\frac{-(t - t_{on})}{T_1}})$$



Spin-Lattice Relaxation Time

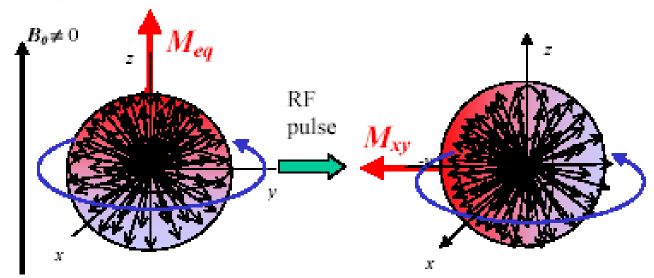
 $R_1 = 1/T_1$ [Hz] longitudinal relaxation rate constant

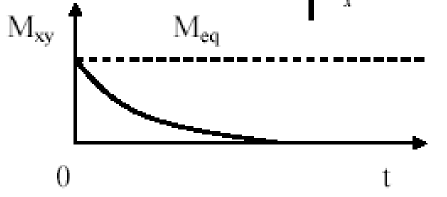
 T_1 [s] longitudinal relaxation time

spin-lattice relaxation time

enthalpy

Transverse Magnetisation





Spin coherence

$$\frac{d}{dt}M_{xy} = -R_2 M_{xy} = \frac{-1}{T_2} M_{xy} M_{xy}(t) = M_{eq} e^{\frac{-t}{T_2}}$$

Spin-Spin Relaxation Time

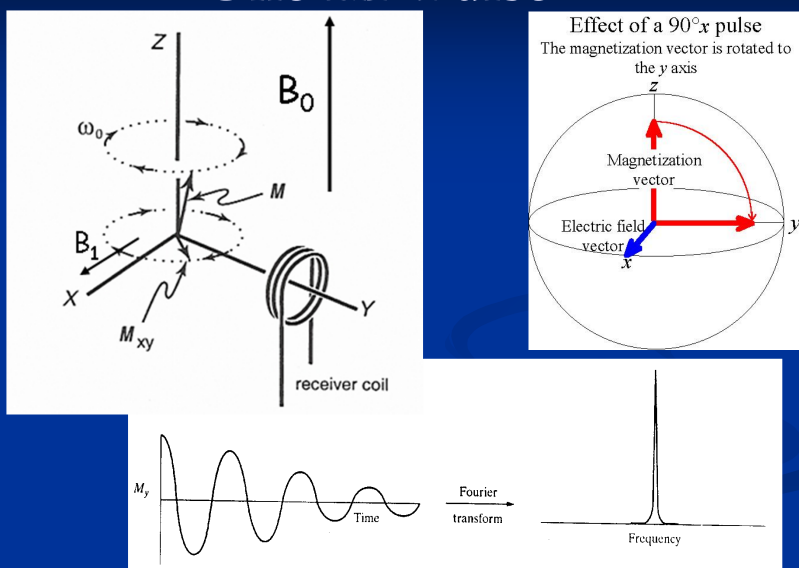
 $R_2 = 1/T_2$ [Hz] transverse relaxation rate constant

 T_2 [s] transverse relaxation time constant

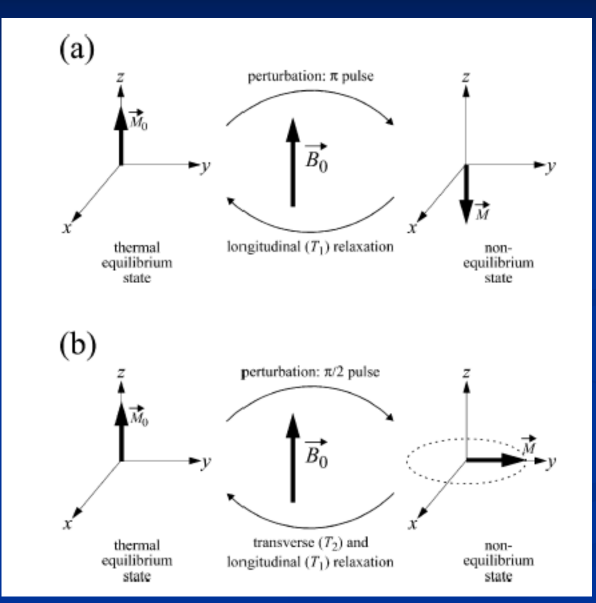
spin-spin relaxation time

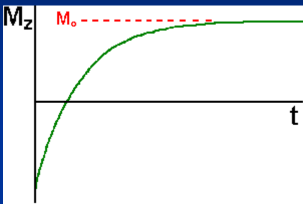
entropy

One RF Pulse

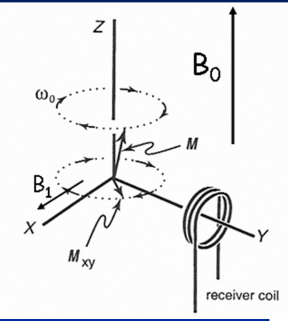


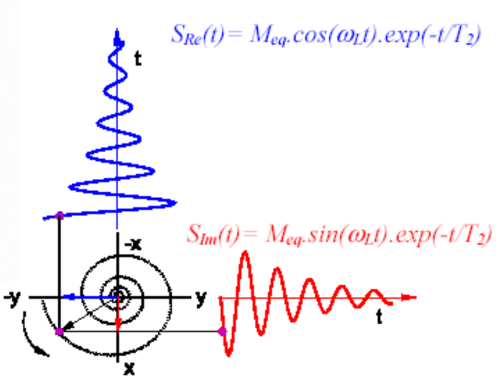
Relaxation





Free Induction Decay FID





$$S(t) = S_{Re}(t) + iS_{Im}(t)$$

=
$$[cos(\omega_L t) + isin(\omega_L t)] exp(-t/T_2)$$

=
$$exp(i\omega_L t)exp(-t/T_2) = exp[-(1/T_2 - i\omega_L)t]$$

