Spin Lattice Relaxation in Solution and Summary of Relaxation Mechanisms

Gareth R. Eaton and Sandra S. Eaton
Department of Chemistry and Biochemistry
University of Denver

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What can you learn from measuring $T_1$ and $T_2$ in solution?

• Dynamics of molecular tumbling
• Oximetry
• Mechanisms of relaxation
  • Spin rotation
  • Modulation of $g$ and/or $A$ anisotropy
  • Modulation of zero-field splitting
Background Information

Early work by CW saturation on Fremy's salt gave $T_1 = T_2 = 0.5 \, \mu s$.
Early work by saturation recovery, mostly of semiquinones by Rengen and later studies by Hyde and co-workers on nitroxy1 radicals also yielded values in the $\mu s$ range.
The rest of the early work on relaxation came largely from low temperature solid state physics.
There developed a dichotomy – solid state effects vs. effects relevant to the fast motional limit.
We have a research program that seeks to find the most important contributions to relaxation in the intermediate region where many EPR studies are performed.
Experimental Methods

In principle, any of the methods discussed for solids could be applied in solution. However, for many samples $T_2$ in fluid solution is so short that it is difficult to perform spin echo experiments. As a result, measurements of relaxation times in fluid solution typically are done by the following methods.

Saturation recovery curves can be measured even when $T_2$ is too short for spin echo formation.

Power saturation curves can be measured on a CW spectrometer. Disadvantages are that a) the saturation behavior depends on the product $T_1T_2$ and b) the saturation may be strongly impacted by spectral diffusion.

If the CW lineshape is dominated by incomplete motional averaging, $T_2$ can be determined from the linewidth after simulation including any unresolved hyperfine splitting.
Nitroxyl Tumbling Correlation Time

The nitroxyl g and A values along the x, y, and z axes are anisotropic. At X-band, the differences between the resonances along the three axes decrease in the order $m_i = -1 > +1 > 0$. As the molecule tumbles, the smaller splitting for $m_i = 0$ is averaged more effectively than the larger splittings, which causes differences in the linewidths of the three hyperfine lines.

Nitroxyl Lineshapes

As the tumbling correlation time decreases, the extent of averaging of anisotropic features increases and the spectrum approaches the 3-line signal that is characteristic of rapid tumbling.

In the motional narrowing region, the dependence of the width of an individual hyperfine line on the nuclear spin state \( m_I \) can be expressed as

\[
\Delta B(m_I) = A + B m_I + C m_I^2
\]

The example on the following page assumes isotropic tumbling.
Calculation of Tumbling Correlation Time

The parameters $B$ and $C$ are related to peak-to-peak amplitudes, $I(m_l)$ by:

\[ B = \frac{1}{2} \left[ \sqrt{I(0)} - \sqrt{I(1)} \right] \quad \text{and} \quad C = \frac{1}{2} \left[ \sqrt{I(0)} + \sqrt{I(1)} - 2 \right] \]

The high-field line has $m_l = -1$.

Tumbling correlation times are calculated from $B$ and $C$ using

\[ \tau = B \left[ \left( \frac{2h}{\sqrt{3} g_o \beta} \right) \left( \frac{4B_o}{15} \right) (b\Delta\gamma) \left( \frac{1}{\Delta B_o} \right) \right]^{-1} \quad \text{and} \quad \tau = C \left[ \left( \frac{2h}{\sqrt{3} g_o \beta} \right) \left( \frac{b^2}{8} \right) \left( \frac{1}{\Delta B_o} \right) \right]^{-1} \]

Where

\[ g_o = \frac{1}{3}(g_x + g_y + g_z) \quad \quad B_o = \frac{\hbar \omega}{g_o \beta} \]

\[ \Delta\gamma = \frac{\beta(g_z - 0.5(g_x + g_y))}{\hbar} \quad \quad b = \frac{2}{3}(A_z - 0.5(A_x + A_y)) \]

$\Delta B_o$ is the peak-to-peak width of the center line.

Hyperfine values (A) are in radians/s

The calculation assumes isotropic tumbling.


Sample Calculation

4-OH-TEMPO (tempol) in 9:1 glycerol:water

g_x = 2.0103, g_y = 2.0067, g_z = 2.0032

A_x = 2\pi \times 18 \times 10^6, A_y = 2\pi \times 22.5 \times 10^6, A_z = 2\pi \times 103 \times 10^6 \text{ rad/s}

I(+1) = 13.5, I(0) = 16.4, I(-1) = 3.4 (arbitrary units)

\Delta B_o = 3.52 \text{ Gauss} \quad \beta = 9.274 \times 10^{-21} \text{ erg/G}

\nu = 9.2449 \times 10^9 \text{ s}^{-1} \quad h = 6.626 \times 10^{-27} \text{ erg s}

\tau = 2.1 \times 10^{-9} \text{ s from B or } \tau = 2.3 \times 10^{-9} \text{ s from C}

The disagreement is an indication of the approximate nature of this calculation. State-of-the-art calculations incorporate detailed physical models.
Slower Tumbling

When tumbling is slower than that required for motional narrowing or tumbling is anisotropic, determination of the tumbling correlation time is best done by lineshape simulation. The results are model dependent.

When a spin label is attached to a protein, rotation of the probe with respect to the protein surface may not be possible through a full 360° and may be highly anisotropic. Freed and his colleagues have modeled these spectra in terms of microscopic order/macroscopic disorder (MOMD).

Extensive discussion of these cases can be found in:

Oximetry

Collisions with paramagnetic oxygen molecules decrease $T_{1e}$ and $T_{2e}$. This is analogous to the use of paramagnetic metals as NMR relaxation enhancement reagents. For narrow-line EPR signals the decrease in $T_{2e}$ causes line broadening, which can be avoided by degassing the sample.

The broadening of narrow-line EPR signals by oxygen can be used to monitor oxygen concentration, which is called oximetry, and is being developed as an in vivo monitoring technique.

The oxygen transport parameter, $W$, is defined as

$$W = T_1^{-1}(\text{air}) - T_1^{-1}(\text{N}_2)$$

and is proportional to the local diffusion coefficient of oxygen.


Oxygen Accessibility

Accessibility of nitroxyl spin labels to collisions with oxygen can be used to distinguish between buried and surface exposed sites on a protein. Measurements could be done by time domain methods, but so far have been done by CW saturation. The accessibility parameter $\Pi$ is defined as:

$$\Pi(O_2) = \frac{\Delta P_{1/2}(O_2)}{P_{1/2}(DPPH)} \frac{\Delta H_0(DPPH)}{\Delta H_o}$$

$P_{1/2}$ is the microwave power that causes the amplitude of an EPR signal to be half of that predicted in the absence of power saturation.

$\Delta P_{1/2}(O_2) = P_{1/2}(O_2) - P_{1/2}(N_2)$

$\Delta H_0(DPPH)$ is the peak-to-peak linewidth of crystalline DPPH (2,2-diphenyl-1-picrylhydrazyl).

$\Delta H_o$ is the peak-to-peak linewidth of the central nitroxyl line, which is a qualitative indicator of mobility. As motion slows, $\Delta H_o$ increases.

Oxygen accessibility and probe mobility were measured as a function of sequence number for spin labels attached to T4 lysozyme (T4L) and cellular retinol binding protein (CRBP). The correlation between the two parameters indicates that the most mobile sites are also the most oxygen accessible. The repeat period of about 3.6 for T4L is consistent with the $\alpha$-helical structure of this segment of the protein.
Molecular Tumbling Induces Relaxation

Molecular tumbling induces relaxation in $S=1/2$ systems by

- spin rotation
- modulation of $g$ anisotropy
- modulation of hyperfine anisotropy

Each of these contributions has a characteristic dependence on tumbling correlation time and microwave frequency. Very few species have been studied as a function of temperature or tumbling correlation time, so it is not known which terms dominate for various classes of radicals.

$$\frac{1}{T_{1e}} = \frac{A_{SR}}{\tau_R} + (A_g + A_a) \frac{\tau_R}{1 + \omega_e^2 \tau_R^2}$$

$\tau_R$ is the tumbling correlation time

$A_{SR}$ is the coefficient for spin rotation and is proportional to $(g-g_e)^2$

$A_g$ is the coefficient for modulation of $g$ anisotropy, which is proportional to $g$ anisotropy squared, and to $\omega_e^2$

$A_a$ is the coefficient for modulation of hyperfine anisotropy, which is proportional to hyperfine anisotropy squared, and is independent of $\omega_e$


Semiquinones

Viscosity, $\eta$, was varied by changing the composition of ethanol:glycerol mixtures at constant temperature. The tumbling correlation time, $\tau$, is expected to be linearly related to $\eta$ so the linear dependence of $1/T_{1e}$ on $1/\eta$ is consistent with spin-rotation. $g$-values are further from 2.0023 for $X=\text{Br}$ than for $X=\text{Cl}$, so the faster relaxation for $X=\text{Br}$ than for $X=\text{Cl}$ also is consistent with spin rotation.

Power saturation curves demonstrated that relaxation was independent of microwave frequency, which is not consistent with modulation of $g$ and/or $A$ anisotropy.

Triarylmethyl Radicals

For these radicals g-values are close to 2, and the hyperfine coupling is weak. As a result, spin rotation and modulation of g and/or A anisotropy are not expected to be effective relaxation mechanisms.

Values of $T_{1e}$ in mixtures of water:glycerol showed little dependence on viscosity or on microwave frequency between 250 MHz and 9.5 GHz, which indicates that spin rotation and modulation of g and A anisotropy do not dominate the relaxation, consistent with expectation.

Values of $T_{1e}$ measured by saturation recovery or inversion recovery were the same, within experimental uncertainty, which indicated the absence of spectral diffusion processes. The 1:1 water:glycerol glass melts at 200 K, but there is no change in the slope of the plot of $1/T_{1e}$ vs. $T$, consistent with the room temperature measurements that indicated no dependence on tumbling correlation time.

The temperature dependence is consistent with a Raman process at low temperature plus a local vibrational mode at higher temperature. The energy for the local mode is consistent with that for the C-S stretch.
Nitroxy Radicals – $T_m$

Values of $T_m$ for tempone in 85% glycerol:15% water obtained by (O) two-pulse spin echo data (Δ) 2D- ESE contour plots
The three lines show the calculated values based on jump diffusion (___), free diffusion (_ _) and Brownian diffusion (_ . _ . ).

In the intermediate tumbling regime $T_m$ for a nitroxy radical is too short to measure by current spin echo techniques.

This is a special case of atomic spectroscopy in condensed phase. The EPR lines are very narrow because there are few neighboring nuclei with a spin. The N ground state is $^4S_{3/2}$ and the hyperfine splitting is isotropic. $T_1$ of N@C$_{60}$ was about 200 $\mu$s in polycrystalline C$_{60}$ at room temperature. In solution the central line exhibited an ESE decay constant of 120 $\mu$s, and $T_1$ was found to be equal to $T_2$. Demonstration that the relaxation of N@C$_{60}$ in solution was a thermally-activated process, attributed to collisional modulation of ZFS for the $S=3/2$ center, was provided by the observation that $T_{1e}$ at 95 GHz was four times longer than at 9.5 GHz. The data fit the following expression, which was derived for $S = 3/2$ ($l = 1,2$):

$$\frac{1}{T_1(l)} = \frac{12}{15} D^2 \left(1 + \frac{\eta^2}{3}\right) \frac{\tau}{1 + (\omega \tau)^2}$$

where D is the ZFS parameter, $\eta$ is the asymmetry parameter, and $\tau$ is the time of the fluctuating process.

Summary of Temperature Dependence

Temperature dependence of relaxation rates observed when $T_m$ is dominated by:

A  nuclear spins in the surroundings
B  collapse of couplings to methyl groups or rapidly relaxing spin
C  averaging of g- or A-anisotropy

$T_1$ is dominated by:

D  direct process and/or high spin concentration
E  Raman or Orbach process
F  interaction with a more rapidly-relaxing electron spin
G  local mode, thermally-activated process, modulating spin-orbit coupling or ZFS
Contributions to $T_1$

• vibrations anisotropically modulate spin-orbit coupling, and mix spin and orbital angular momentum
• hence, for axial symmetry, $T_1$ is shortest in the xy plane, and longest along z
• vibrations of the molecule exchange energy with the lattice
• collisions of the molecule with solvent (including in frozen solution) excite vibrations of the molecule
• at ca. 100 K in a "frozen" solution a molecule undergoes low-amplitude molecular motions many times during $T_1$
• in a "rigid" lattice energy can be exchanged with the lattice via a single resonant phonon – direct process two phonons – Raman process excited state electronic energy level within the phonon bath – Orbach process excited state vibrational energy level within the phonon bath – Murphy process local vibrational mode special vibration/phonon interactions also are possible
• temperature dependence can be complicated, and includes linear ($T_1 \propto T^{-1}$) to $T_1 \propto T^{-9}$, but is commonly $T_1 \propto T^{-n}$ with $n = 2$ to 3 from ca. 10 K to the melting point of the solvent.
• spin-spin interaction with a faster-relaxing electron spin
Contributions to $T_m$

• $T_2 = \text{electron spin-spin relaxation}$
• Instantaneous diffusion. The second pulse flips a neighboring spin.
  The process has larger impact for
  higher spin concentrations, narrower EPR signals, and
  larger microwave $B_1$
• Intramolecular dynamic processes
  small amplitude motion
  rotation of methyl groups to which the unpaired electron is
  coupled
  relaxation of spin-coupled systems
• Averaging of $g$ and $A$ anisotropy by molecular tumbling
• Nuclear spins in the surroundings
Summary of Relaxation Mechanisms

The spin angular momentum of the A spins is mixed with other angular momenta by spin-orbit coupling or spin-rotation coupling. The time-dependence of the mixing of angular momenta, occurs by rotational motion, motional modulation of dipolar coupling, mixing of excited states (e.g. by conformational changes), thermal motions modulating zero-field splitting.

This mixing of angular momenta is thermally activated, and provides a mechanism for coupling the spin angular momenta to the thermal energy of the lattice.
Relaxation Mechanisms

- Microwaves
- Resonant Spins (A spins)
- Longitudinal
- Enthalpy
- T1
- Transverse
- Entropy
- T2, Tm
- Spin diffusion
- Cross relaxation
- Heisenberg exchange
- T1, T2
- Spin-orbit
- ZFS
- Local modes
- Spin-rotation
- Spin diffusion
- Instantaneous diffusion
- Rotations
- Heisenberg exchange
- B spins
- Other Angular Momenta
- Direct
- Raman and Orbach
- Lattice Thermal Motions
- Bath
Further Reading on Electron Spin Relaxation Times


