Mechanical detection of nuclear spin relaxation in a micron-size crystal

O. Klein\textsuperscript{1} \textsuperscript{a}, V. V. Naletov\textsuperscript{2}, and H. Alloul\textsuperscript{1} \textsuperscript{b}

\textsuperscript{1} Laboratoire des Solides Irradiés, Ecole Polytechnique, Palaiseau F-91128
\textsuperscript{2} Physics Department, Kazan State University, Kazan 420008 Russia

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Abstract. A room temperature nuclear magnetic resonance force microscope (MRFM), fitted in a 1Tesla electromagnet, is used to measure the nuclear spin relaxation of \textsuperscript{1}H in a micron-size (70ng) crystal of ammonium sulfate. NMR sequences, combining both pulsed and continuous wave r.f. fields, have allowed us to measure mechanically $T_2$ and $T_1$, the transverse and longitudinal spin relaxation times. Because two spin species with different $T_1$ values are measured in our 7\,$\mu$m thick crystal, magnetic resonance imaging of their spatial distribution inside the sample section are performed. To understand quantitatively the measured signal, we carefully study the influence of the spin-lattice relaxation and the non-adiabaticity of the c.w. sequence on the intensity and time dependence of the detected signal.

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1 Introduction

For a long time research groups have looked for new ways of detecting electronic or nuclear paramagnetic resonance with better sensitivity. A review of different proposed methods can be found in the introduction of Abragam's book [1]. In two seminal papers [2,3], Sidles recognized the advantage of coupling the spin system to a mechanical oscillator for magnetic resonance imaging. In this technique, the force signal is proportional to the magnetic field gradient [4], which, in extremely inhomogeneous field, should allow high spatial resolution. The new technique is referred to as magnetic resonance force microscopy (MRFM) [5].

The first magnetic resonance force signal was detected by Rugar et al. in 1992 while exciting electron spin resonance (eMRFM) in a 30ng crystal of diphenylpicrylhydrazil [6]. Two years later, Rugar et al. reported the mechanical detection of \textsuperscript{1}H (protons) nuclear magnetic resonance (nMRFM) in 12ng of ammonium nitrate [7]. These two pioneering experiments demonstrate that a micro-fabricated cantilever, identical to the ones developed for atomic force microscopy, can detect the magnetic moment of a micron-size sample. In the case of nuclear magnetic resonance (NMR) [7], the achieved sensitivity of $10^{13}$ spin, at room temperature and in a field of 2.4T, represents a substantial improvement over the standard coil detection.

Significant progress was made in the past few years. In 1996, Zhang et al. mechanically detected the ferromagnetic resonance (fMRFM) of yttrium iron garnet [8]. Imaging experiments with eMRFM [9,10], nMRFM [11,12] and fMRFM [13] were performed. A magnetic resonance torque signal in a homogeneous magnetic field [14] was also detected. Improvement of the force sensitivity by operating at low temperature [15–17] was demonstrated. Force maps of the sample were obtained with the magnetic probe placed on the mechanical resonator in eMRFM [18] and fMFRM [13]. The highest sensitivity reported so far is around 200 electron spin in a 1Hz bandwidth. The result was obtained by operating an eMRFM at 77K in a very large magnetic field gradient [19]. In 1996, Wago et al. demonstrated that a pulse sequence combined with fast adiabatic passages can allow to measure the nuclear spin-lattice relaxation time of \textsuperscript{19}F in calcium fluoride at low temperatures [15]. The same method was used to measure the longitudinal spin relaxation of \textsuperscript{1}H in ammonium sulfate at room temperature and normal pressure [20,21]. Recent eMRFM work in vitreous silica at 5K showed that the same principles can be also applied to study electron spin dynamics of E\textsuperscript{′} centers with long $T_1$ [17].

In this paper, we report the first measurements of both the transverse and longitudinal nuclear spin dynamics of \textsuperscript{1}H using mechanical detection. A very thin sample is used to analyze if new phenomena might be specific to small sizes. Our instrument is a simple home-built...
MRFM located inside a 1 Tesla electromagnet. The mechanical motion of the cantilever is monitored by a laser beam deflection system. The sample is a 7 μm thick crystal of (NH₄)₂SO₄. Two spin-lattice relaxation times are observed, T₁ ≈ 0.4 s and T₂ ≈ 5 s. The later value corresponds to the T₁ reported in the literature for this compound [22,23]. The short relaxation, however, might be due to water contamination inside the crystal during its contact with air. These same two relaxation rates are also measured by conventional NMR in powder samples with particles of dimensions smaller than 50 μm.

After introducing in Section 2 the measurement technique employed in this study, we will present in Section 3 our results on the transverse and longitudinal spin relaxation properties of (NH₄)₂SO₄. This will be followed in Section 4 by a more detailed analysis of the time dependence and magnitude of the force signal in order to quantify the properties of spin with short and long T₁ and to determine the effect of the non-adiabaticity of the sequence in the measured signal. Finally a model to describe our experimental data will be proposed.

2 Measurement of the force signal

The setup is schematically represented in Fig.1. The experiment is performed at room temperature inside a vacuum cell (10⁻⁵ torr) constantly connected to the inlet of a rotary pump. The instrument [24] fits between the poles of an iron core electromagnet which produces a static magnetic field B_{ext} = 1 T. Near the uniform field one adds a second inhomogeneous field with axial symmetry produced by a magnetized iron bar 8 mm in length and 1.9 mm in diameter. The polarization field experienced by the spin is B₀ = B_{ext} + B_{pol}, with B₀ = B_{0} · k and B_{pol} = B_{cyl} · k. Near the symmetry axis, the instantaneous magnetic force acting on the sample is given by the expression [25]:

\[ F(t) = \int_{V_s} M_z(r,t) \frac{\partial B_{cyl}}{\partial z} dV. \]  

Here M_z is the z component of the bulk magnetization and V_s is the volume of the sample. For small sample size, we make the approximation that the field gradient g = \( \partial B_{cyl}/\partial z \) is uniform over V_s. A new length variable \( \zeta = B_0(r)/g \) is defined so that a plane of constant \( \zeta \) maps onto a surface (actually a paraboloid) of constant polarization field which also corresponds to a sheet where the spin have the same motion. A paraboloid of fixed \( \zeta \) value, however, shifts axially away from the iron cylinder when B_{ext} increases. In this experiment, the sample is placed 0.70 mm above the iron cylinder and centered on the cylinder axis. At this distance, the calculated axial field gradient is g = -470 T/m (see appendix A).

The mechanical force detection is obtained by measuring the elastic deformation along the z axis of a micro-fabricated cantilever on which the sample is attached. In this orientation, the probe is sensitive to the longitudinal component of the nuclear magnetization in contrast with a standard coil detection. The cantilever equation of motion is represented by a damped harmonic oscillator with a single degree of freedom. The measurement technique uses the optical deflection of a 4 μW HeNe laser beam which reflects off the rear side of the cantilever onto a position-sensitive detector.

Our test compound is (NH₄)₂SO₄. This non-magnetic insulator has a high proton density \( d = 6.4 \times 10^{22} \text{H/cm}^3 \) and is in its paraelectric state above 223 K. NMR measurements of the \(^1\text{H}\) spin-lattice relaxation time at 300 K in our powder [26] give T₁ ≈ 5 s along the static field [27]. The \(^1\text{H}\) linewidth is 5 G and the second moment is M₂ = 4G² at 295K [28]. Our sample is a crystal cleaved to a platelet aspect ratio and glued with epoxy on the end of a commercial Si₃N₄ amorphous cantilever of spring constant k = 0.008 N/m as can been seen in Fig.2. After completing the assembly, the cantilever resonance frequency drops from 5.8 kHz to 1.4 kHz due to the sample mass [29]. The quality factor of the loaded cantilever is \( Q \approx 4000 \) in vacuum. From the electron microscopy images (Fig.2), the sample dimensions are approximately 100 × 50 × 7 μm³ with the smallest length (the thickness) oriented along the axial field. This represents a volume \( V_s = 3.5 \times 10^{-8} \text{cm}^3 \) or a mass m = 70 ng and corresponds to \( N \approx 10^{15} \) protons. The temperature of the cantilever holder is stabilized around +27°C during the measurement [30]. The nuclear magnetization at thermal equilib-
rium is expressed by the Curie law \( M_0 = (d\mu_n^2 B_0/k_B T) \mathbf{k} \), with \( \mu_n = 1.4 \times 10^{-26} \text{J/T} \) the proton magnetic moment, \( T = 300 \text{K} \) and \( B_0 = 1.3 \text{T} \) the polarization field. This gives a magnetic moment \( M_0 V_x = 2.3 \times 10^{-16} \text{J/T} \).

In order to increase the sensitivity, \( M_z \) is modulated at a frequency \( \omega_m \) close to \( \omega_c \), the frequency of the fundamental flexure mode of the cantilever. At the moment the optimal configuration uses cantilevers that have mechanical resonance frequencies in the audio range and Larmor frequencies \( \omega_L \) which are several orders of magnitude larger (radio or microwave frequencies). Only two methods have been used to create an oscillatory force on the cantilever: cyclic saturation and cyclic adiabatic inversion. They are restricted respectively to compounds that have spin-lattice relaxation times \( T_1 \) either shorter or larger than the oscillation period of the cantilever.

In our case, the modulation of \( M \) is generated by a continuous-wave (c.w.) sequence that consists of periodic adiabatic fast passages [1]. The radio-frequency (r.f.) source is a 35-75MHz Voltage Controlled Oscillator (VCO). The r.f. output field is amplified up to 7W and fed into an impedance matched resonating circuit \( (Q_{rf} \approx 100) \) tuned to a fixed frequency, 54.7MHz. A small coil (3 turns, 0.8mm in diameter) is in series with the tank circuit. The sample is 0.5mm away from this antenna. The nuclear gyromagnetic ratio \( \gamma = 1.4 \times 10^8 \text{s}^{-1} \text{T}^{-1} \) in vacuum. The estimated sample volume is \( 3.5 \times 10^{-7} \text{cm}^3 \).

The surface of constant \( \zeta = \zeta_0 \) is called the resonant sheet. The sinusoidal frequency modulation is started at a time \( t = 0 \). In a transformation to a rotating coordinate system with an instantaneous angular velocity \( \omega(t) \mathbf{k} \), the apparent magnetic field is:

\[
B_z(\zeta, t) = B_3 + \left\{ \frac{g \zeta - \omega(t)}{\gamma} \right\} \mathbf{k}.
\]

\( \theta \) is defined as the polar angle made by the apparent field with the external field. The magnetization, however, progresses about the direction \( B_e + \theta/\gamma \mathbf{j} \), with \( \theta = \partial \theta/\partial t \) (see appendix B). A parameter for non-adiabaticity is defined with \( \tan \alpha = \theta/\gamma | B_e | \) the angle between the two vectors. Provided that the adiabatic condition \( \alpha \ll 1 \) is satisfied, the spin system remains at all times in a state of internal equilibrium and \( M \) is parallel to \( B_e \) as required by Curie’s law. The longitudinal magnetization is:

\[
M_z(\zeta, t) = |M| \cos \theta,
\]

where

\[
\cos \theta = \frac{g \zeta - \omega(t)/\gamma}{\sqrt{(g \zeta - \omega(t)/\gamma)^2 + B_3^2}}.
\]

For free spin, \(|M|\) is a constant of the motion [1]. This is no longer true in condensed matter because of spin-lattice relaxation. In our sample, however, the magnetization decay is slow compared to the modulation period. Under our measurement protocol, an extra defocusing originates from the lack of adiabaticity of the modulation. In a first step, these effects are neglected and they will be considered in a more detailed analysis deferred to a later section (see also the appendix B). At time \( t = 0 \), \( B_3 \) is assumed to be turned on adiabatically with the sample initially in thermal equilibrium. In this case the norm \( M \) reflects the state of the longitudinal magnetization immediately before the force measurement. During the c.w. sequence, the oscillatory movement of \( M_z(t) \) comes from the \( \cos \theta \) factor. The value is expanded in time series \( \cos \theta \approx a_0 + a_{1} \sin(\omega_m t) \) [31] with \( a_{1} \) the first harmonic Fourier component [11] (higher harmonics have a negligible effect on the motion of the cantilever). Because of the large field inhomogeneities, the amplitude of oscillation depends on the location inside the sample. The resonant sheet, which is the paraboloid of constant \( \zeta_0 \), corresponds to the surface of maximum amplitude of oscillation. The spatial dependence of \( a_{1}(\zeta) \) is the sensitivity profile. \( \Gamma \) is the half width at half maximum of this bell-shaped curve. \( \Gamma \) has the units of a distance and it defines the thickness of the slice probed. The amplitude of \( \Gamma \) depends on both \( \Omega \) and \( B_3 \) [32]. The induced vibration is synchronously amplified by a lock-in technique through a single-pole low-pass linear filter of time constant \( \tau_1 \). For \( \omega_m = \omega_c \), the lock-in signal grows exponentially (an exact expression will be given in equation (11)) to the asymptotic amplitude:

\[
A_0 = \frac{1}{\sqrt{2}} \frac{Q g}{k} \int_{\zeta_0} M_0 a_{1}(\zeta) d\zeta.
\]

In conclusion, the maximum amplitude of vibration achieved by the cantilever is proportional to the longitudinal
At the start, the amplitude of $B$ is set to 27s ($\omega_1$). The time delay between force measurements is the lock-in output $A$. The crystal is irradiated for 3s by a r.f. field of $\tau = 10G$ (upper panel). The bottom panel shows the frequency waveform applied to the VCO around $\omega_1 = 54.7$MHz. The amplitude of the frequency modulation is $\Omega/2\pi = 150$kHz. The width of the curve is the digitalization noise of the oscilloscope (not the phase noise).

Fig. 3. (a) Vibration amplitude of the cantilever measured by the lock-in for a (NH$_4$)$_2$SO$_4$ crystal containing $10^{15}$ protons at 300K in $B_{ext} = 0.9425T$. The trace corresponds to a single shot experiment with no averaging. The lock-in time constant is $\tau = 0.3$s. (b) Details of the start and end of the c.w. sequence. The crystal is irradiated for 3s by a r.f. field of $B_1 = 10G$ (upper panel). The bottom panel shows the frequency waveform applied to the VCO around $\omega_1/2\pi = 54.7$MHz. The amplitude of the frequency modulation is $\Omega/2\pi = 150$kHz. The width of the curve is the digitalization noise of the oscilloscope (not the phase noise).

magnetic moment inside the probed slice at the beginning of the c.w. sequence. In the ideal case of a uniform inversion of all spin inside the sample, the asymptotic amplitude would be $A_{tot} = Q\mu_0 M_0 V_c/(k\sqrt{2})$.

In the upper panel of Fig.3 the time dependence of the lock-in output $A(t)$ is shown when the c.w. sequence is applied. The time delay between force measurements is set to 27s ($> 5T_1$) to ensure a steady state magnetization close to the thermal equilibrium value. The lock-in time constant is $\tau = 0.3$s which corresponds to an output noise of 4A. The bottom panel of Fig.3 displays the time dependence of $B_1$ and $\omega$ at the beginning and end of the c.w. sequence. The amplitude of $B_1$ is turned on from 0 to 10G in 5ms when the frequency is well off-resonance, i.e. $400$kHz below $\omega_1/2\pi$. The frequency is then ramped to resonance in 7ms. Finally, the frequency modulation of the r.f. field is applied for 3s with a deviation $\Omega/2\pi = 150$kHz. For these settings, the calculated value of $\Gamma = 7\mu$m is comparable to the sample thickness.

Since the r.f. tank circuit is tuned to a fixed frequency, the resonance is found by sweeping the external field $B_{ext}$. There are no spurious vibrations of the cantilever induced by the r.f. fields when $B_{ext}$ is outside the resonance range. Fig.3a shows the amplitude of the lock-in signal achieved in a one shot experiment at the resonance maximum, $B_{ext} = 0.9425T$. The maximum vibration amplitude is around 40A which corresponds to a signal to noise ratio of 20dB.

3 Relaxation measurements

In this section the nuclear spin dynamics of our sample is measured by applying a series of r.f. pulses before the c.w. sequence described above.

In order to calibrate the strength of the r.f. field, a r.f. pulse of duration $\tau_p$ is applied, with an amplitude $B_1$, 13ms before the c.w. sequence. During this pulse, $M_0$ rotates about $B_p$ through an angle $\varphi$. The angle obtained at the end of the pulse is $\varphi = \gamma |B_p| \tau_p$ [15,17]. Within a few milli-seconds after the pulse, the nutated magnetization vector decays to its longitudinal component which then determines the amplitude of the maximum vibrations achieved by the cantilever during the force measurement. $B_1$ is set at maximum power, i.e. 6.4W, for the pulse. The c.w. sequence uses a 2.9W r.f. field. Fig.4 shows the lock-in output averaged over a 1s time interval around its peak
amplitude. For spin that are at $\zeta_0$, the signal is proportional to $\cos \varphi$. The data are fitted by the functional form $\exp(-\tau_p/\tau) \cos(\gamma B_1 \tau_p) + b$. The period gives a calibration of the r.f. field strength at the sample location and we get $B_1 = 15G$ during the pulse. The other fitting parameters are $\tau = 43 \pm 6\mu s$ and a positive offset $b = 3 \pm 0.2\mu A$. The values of these last two parameters depend strongly on $B_1$. The positive offset $b$ is mainly due to the non-uniform field inside the sample [33]. For $^1H$ away from $\zeta_0$, the direction of $\mathbf{B}_0$ is not exactly perpendicular to $\mathbf{k}$ and only a partial inversion of the $z$ component is obtained when $\varphi = \pi$. The decay of the magnetic moment fitted by $\tau$ is due to field inhomogeneity which causes a dephasing of the magnetization in the transverse plane [34].

To study the transverse magnetization decay of $^1H$ [17] a sequence of 3 pulses is used. A $\pi/2$ pulse is applied to the spin system, so that the magnetization at $\zeta_0$ is rotated to the transverse plane. After a fixed delay $\tau_a$, a $\pi$ pulse is applied to form a spin echo. Shortly after, a $\pi/2$ pulse takes an instant snap-shot of the transverse magnetization by rotating it along $\mathbf{k}$ and the frozen component is measured with the c.w. sequence. Varying the time delay $\tau_b$ between the last two pulses reconstructs the transient shape of the spin echo. Using the same settings as the earlier measurement, the widths of the $\pi/2$ and $\pi$ pulse are set to $3.8\mu s$ and $7.6\mu s$ respectively. The delay between the center of the first two pulses is $\tau_a = 17\mu s$. In Fig.5, the lock-in peak (again averaged over 1s around its maximum) is shown as a function of $\tau_a + \tau_b$. As expected for a spin echo, the reconstructed transverse magnetization becomes refocused at a time $2\tau_a$.

With increasing spacing $\tau_a$ between pulses, the size of the spin echo signal decreases due to spin-spin relaxation. Using the same sequence as above, Fig.6 is a plot of the spin echo amplitude measured as a function of the time $2\tau_a$. The measured values are plotted on a $x^2 - \log(y)$ scale and one finds that the data follow the relationship $\exp\{-t/2T_2\}$ with $T_2 = 39 \pm 1\mu s$. With the inferred $T_2$, the shape of the echo in Fig.5 can be calculated taking into account the dipolar linewidth of the protons in our compound [28] and the spatial dependence $a_1(\zeta)$. The solid line in Fig.5 is the best fit obtained for a sample thickness of $6.5\mu m$ which is in good agreement with the value obtained on the image.

The longitudinal magnetization recovery is now measured after a saturation comb [35]. This protocol puts efficiently inhomogeneous spin systems in a well defined uniform state outside thermal equilibrium. The saturation comb is composed of three $\pi/2$ pulses spaced by $100\mu s$. The c.w. sequence is applied at a variable delay (13ms $< t < 20s$) after the comb. In order to obtain an intrinsic measurement of the relaxation, it is important to ensure that the sensitivity profile $a_1(\zeta)$ is exclusively included inside the sample section, otherwise a partial re-polarization of the magnetization occurs during the measurement cycle [17]. For our settings, $\zeta_0$ is set exactly at the middle of the sample and $\Gamma = 2.4\mu m$ is smaller than the sample thickness. As before, the value plotted is the lock-in output averaged over a 1s time interval around its maximum. No signals are detected when $t = 13ms$. On Fig.7, two relaxation times in the recovery process are clearly observed. The results are fitted with a double exponential $\phi_x \{1 - \exp(-t/T_{1x})\} + (1 - \phi_x) \{1 - \exp(-t/T_{11})\}$ which gives $\phi_x = 49 \pm 2\%$, $T_{1x} = 0.35 \pm 0.03s$ and $T_{11} = 5.4 \pm 0.5s$. The value $\phi_x$ does not correspond directly to the proportion of spin that have a short relaxation ($n_x$) since the
factor between $M_z$ and the lock-in amplitude is also a function of $T_1$.

Neutron diffraction studies [36] of the (NH$_4$)$_2$SO$_4$ crystal structure show that there are two NH$_4^+$ sites at room temperature surrounded respectively by five and six SO$_4^{2-}$ ions. The protons of the two inequivalent ammonium ions are coupled via dipole-dipole interactions and the measured spin-lattice relaxation rate at 300K is an averaged value of the $T_1^{-1}$. In a variable temperature NMR measurements, O’Reilly and Tsang [23] observe a single exponential $^1$H relaxation process and analyze their $T_1$ results by the reorientation correlation times $\tau_0$ of the two distinctive NH$_4^+$.

At 300K, $1/\tau_0$ should be larger than the Larmor frequency and the rotation should be isotropic which means that $T_1$ should be independent of the orientation between the static field and the crystallographic axis of our sample. We suppose that our observed two $T_1$ processes might be due to the exceptionally small thickness of our crystal (7 $\mu$m).

In order to check the later hypothesis, a conventional NMR measurement was performed by A. Dooglav with a 1T custom spectrometer. The sample consisted of $\sim 1g$ of our sample ground to small particles with dimensions below 50 $\mu$m. In this fine powder sample, a double relaxation process is also observed with the following parameters $n_s = 17 \pm 5\%$, $T_{1s} = 0.37 \pm 0.1s$ and $T_{1l} = 4.7 \pm 0.2s$. This result is in sharp contrast with experiments performed on coarser grains, where only one relaxation is observed with $T_1 = 5.0 \pm 0.2s$. The values of the two relaxation times are equal, within error bars, to the ones measured by MRFM. In addition, measurements were performed on the same 50$\mu$m powder after two weeks of aging in air. It showed a rise of $n_s$ to $26 \pm 3\%$ in the longitudinal magnetization recovery experiment. A standard spin-spin relaxation measurement on this powder seems also to indicate a double time $T_2$ with $n_s = 20 \pm 6\%$, $T_{2s} = 49 \pm 12\mu s$ and $T_{2l} = 79 \pm 1\mu s$.

One corollary issue concerns the spatial distribution of each spin species inside the sample section. To perform this measurement, we record the amplitude of the lock-in signal as a function of $B_{ext}$ for two delays $t$ between the saturating comb and the c.w. sequence. By sweeping $B_{ext}$, the surface $\zeta_0$ is displaced to a different height in the sample. The force signal is then proportional to the density of spin around this location. In order to obtain a local measurement, the thickness $\Gamma$ of the slice probed is reduced by decreasing both $\Omega$ and $B_1$ during the c.w. sequence. The inset of Fig.8 shows the spatial dependence of the transfer function $a_1(\zeta)$ for our settings where $\Gamma$, the half width at half maximum, is 1.9$\mu$m. By changing the delay $t$, the weight $a_\phi$ of one spin species compared to the other can be varied. Qualitatively, the measurement protocol gives more weight to the spin species with short relaxation when the comb is close to the c.w. sequence. The obtained results are shown in Fig.8 for both $t = 0.6s$ (closed circles) and $t = 16s$ (open circles). A first look at the result indicates that a more rounded distribution is obtained for the $t = 0.6s$ data. The measurements, however, collected close to the edge of the sample are skewed by repolariza-
tion processes that modify the shape of the lock-in signal. Inside the bulk of the crystal (0.942T ≤ B_{ext} ≤ 0.944T), there is no clear evidence of a spatial modulation of one spin population compared to the other, e.g. a dip of the signal in the middle of the crystal. This result suggests that, within our resolution, the water contamination is uniform in the thickness. The solid line is a calculation of the expected profile for a parallelepiped sample of dimensions 100 × 50 × 7μm³ within both the free spin and adiabatic approximations. In spite of the idealized model, the t = 16s data (open circles) are well described by the calculated profile except for the high field range. The shoulder at B_{ext} = 0.946T corresponds to the surface of the sample that has been glued with epoxy to the cantilever. We did not attempt to fit this part of the data. The observed step in the signal might be due to the protons in the epoxy. A small roll angle between the sample and the cantilever combined with the particular shape of our crystal is also consistent with the observed effect.

Although the data analyzed here-above ensure that two populations of spin with different NMR properties are present, their actual proportion is not quantitatively determined, as the actual values of the relaxation times influence the magnitude of the measured signal. For a better knowledge of the sensitivity of the technique to the measurement parameters, it is then necessary to perform quantitative analyses.

4 Quantitative measurements

In this section, we shall first calibrate the mechanical response of the cantilever and the mechanical noise. The time dependence A(t) of the lock-in signal is calculated, taking into account relaxation processes and non-adiabatic effects. The experimental responses for different values of B_{1} and Ω are compared with the calculations. This allows us to select an experimental condition for which non-adiabatic effects can be neglected. It is then shown that two relaxation times are indeed required to fit the time dependence of the observed lock-in signals, with values consistent with those obtained from T_{1} data.

The Q of the cantilever is first measured carefully through the noise vibration spectrum of the cantilever loaded with the sample in vacuum when the r.f. power is off. The lock-in time constant is set to τ_{l} = 10s. An audio generator sweeps the lock-in reference around ω_{c}. The plotted value in Fig.9 is the standard deviation of the lock-in signal estimated over 100 × τ_{l} (the mean lock-in signal is zero). During the whole experiment, the temperature stability of the cantilever is better than ±0.01°C which guarantees that ω_{c} does not shift by more than 0.01Hz. Fitting the squared amplitude with a Lorentzian [37], one obtains the cantilever resonance frequency ω_{c}/2π = 1397.77Hz and quality factor Q = 4000 (defined as the ratio of ω_{c} over the full width at half maximum of the power spectrum). Away from resonance, our sensitivity is limited by the noise of the detection electronics. It is several orders of magnitude smaller than the A-scale motion of the cantilever at resonance and therefore it can be neglected.

Near ω_{c}, the cantilever motion consists of white noise amplified by a narrow-bandwidth mechanical resonator [38]. ∆ν_{c} is the one-sided equivalent noise bandwidth (ENBW) of the mechanical resonator ∆ν_{c} = ω_{c}/8Q = 0.27Hz. The noise at the output of the lock-in is this narrow-band noise observed through a RC filter of time constant τ_{l} = 1/RC whose ENBW ∆ν_{l} = 1/4τ_{l} = 0.025Hz. Exactly at resonance, the combined distribution gives an ENBW ∆ν = (1/√{Δν_{c} + 1/Δν_{l}})^{-1} = 0.023 Hz. To convert our data to spectral density, the resonance amplitude in Fig.9 must be divided by √{Δν}. The measured noise spectral density is A = 13A/√Hz. This figure also corresponds to the noise observed in Fig.3 where ∆ν = 0.20Hz. The result has to be compared with the intrinsic correlation function for fluctuations of a Brownian particle harmonically bound to a single degree of freedom oscillator of spring constant k:

\[
\langle AA(t) \rangle = \frac{4Q k B T}{k \omega_{c}},
\]

Taking the square root of the above expression, one gets A_{τ} = 10A/√Hz at T = 300K which is in good agreement with our measured value. In conclusion, our dominant noise comes from the thermal vibration of the cantilever. From this result, one can estimate the smallest force detectable by the instrument in one shot kA/Q = 2×10^{-15}N/√Hz. In order to obtain a quantitative measurement of our force signal when the r.f. field is applied, a more detailed study of the time dependence of the lock-in signal A(t) is needed. The length of the c.w. sequence is increased to 6s compared to Fig.3. Fig.10 is the average of the lock-in signal over 32 sequences using a short lock-in time constant of τ_{l} = 30ms. The striking feature of this plot is...
that the norm of the magnetization $|M|$ decays during the c.w. sequence. We have checked that experimental perturbations such as the phase noise of the r.f. source are negligible in this case [39]. In the rotating frame, the magnetization tends to recover slowly towards a steady state value due to spin-lattice relaxation processes [40]. These relaxation processes are different from the relaxations measured in section 3 which are linked to the time dependence of the magnetization in the absence of r.f. fields. In addition, the lack of adiabaticity quantified by $\alpha \approx \dot{\theta}/\gamma B_0$ produces a precession movement around the locally changing effective field direction. This mistracking corresponds to a magnetization component perpendicular to the instantaneous precession axis $B_0 + \dot{\theta}/\gamma J$ that relaxes due to spin-spin interactions. But in the limit where $\alpha \ll 1$ and $2\pi/\omega_c \ll T_1$, the decrease of $M$ after one cycle $\Delta = M(t+2\pi/\omega_c) - M(t)$ is small. The value is calculated at the lowest order for one spin species (see appendix B).

$$\Delta \approx -M \left\{ \int_0^z \dot{\theta}(t) \left\{ \int_0^t \dot{\theta}(t') \exp \left( -\int_{t'}^t \frac{1}{T_1} dt'' \right) \times \cos \left( \int_{t'}^t \frac{\gamma B_1}{\sin \theta(t'')} dt'' \right) dt' \right\} dt + \frac{z^2}{T_{1z}} \cos \theta(t) + \frac{z^2}{T_{1z}} \theta(t) \right\} + M_0 \int_0^z \frac{\cos \theta(t)}{T_{1z}} dt,$$  

with $T_{1z}$ is the usual $T_1$ in the absence of r.f. field, $T_{1z}$ is the transversal spin-lattice relaxation and $1/T_{1z}^+ = (1/T_{1z} + \cos^2 \theta/T_{1z} + \sin^2 \theta/T_{1z})/2$. It was shown that the relaxation mechanisms in this compound are associated with the time varying field induced by the change in the NH$_4^+$ orientation. For an exponential correlation function with correlation time $\tau_0$, $T_{1z}$ is expressed as a sum of the spectral density of these fluctuating fields $J^{(i)}(\omega) = \tau_0/(1 + \omega^2 \tau_0)$ with an index $i$ that corresponds to the number of net spin flip: $1/T_{1z} = 3/2\gamma^2 \hbar^2 I(1) + 1/2(5/2J^{(1)}(\omega_0) + 1/4J^{(2)}(\omega_0) + 1/4J^{(4)}(2\omega_0))$, with $\omega_c = \gamma B_c$. In the approximation for our compound that $\omega_c \tau_0 \ll 1$, one obtains that $T_{1z}$ is independent of $\theta$ and $T_{1z} = T_{1z}$. Coming back to equation (6), one recognizes that the first integral is due to the finite value of the non-adiabaticity parameter $\alpha$ (see appendix B). The second represents the spin-lattice relaxation $T_{1p}$ in the rotating frame [41]. The last integral is the equilibrium magnetization that corresponds to the spin temperature in the rotating frame. $\tau$ and $M_{eq}$ are defined by rewriting the above expression in the form $\Delta/2\pi = -M/(\omega_c \tau) + M_{eq}/(\omega_c \tau)$. During the c.w. sequence, the oscillatory driving magnetic force is damped and the instantaneous value is given by:

$$F(t) \approx g \sin(\omega_m t) \int V_{xy} a_1(\xi) M(\xi, t) d\xi + \text{constant},$$  

(7)

with

$$M(\xi, t) = \left\{ M_{eq}(\xi) + \left\{ M_0 - M_{eq}(\xi) \right\} \exp \left( -\frac{t}{\tau_m(\xi)} \right) \right\}.$$  

(8)

The integral in equation (7) relaxes approximately according to a single exponential towards its equilibrium value $m_{eq} = \int V_{xy} a_1(\xi) M(\xi) d\xi$ with an apparent characteristic time $\tau_m$. One notes that $m_{eq} = 0$ by symmetry when $\xi_0$ is centered at the middle of the sample $\xi_0 = \xi_m$. The value of $m_{eq}$ is positive for $\xi_0 < \xi_m$ and changes sign for $\xi_0 > \xi_m$ [17]. In the particular case where $m_{eq} = 0$ and $\alpha \ll 1$, then it can be shown that the coefficient $\tau_m$ is bounded between $T_{1z} \leq \tau_m \leq T_{1z}$ [21].

The forced vibrations of an harmonic oscillator are given by the convolution product:

$$a(t) = \beta \int_0^t \frac{F(t')}{k} e^{-\frac{t - t'}{\tau_c}} \sin \left\{ \omega_c(t - t') \right\} \omega_c dt',$$  

with $\beta = \{1 + 1/(4Q^2)\}$ and $1/\tau_c$ the damping constant of the cantilever. In our experiment the external force is $F(t) = F_0 \exp(-t/\tau_m)$ for $m_{eq} = 0$, with $F_0 = kA_0/Q$ and $\tau_m$ the characteristic decay time of the magnetic force. $a(s)$ the Laplace transform of equation (9) is calculated in the complex plane:

$$\frac{k}{F_0} a(s) = \frac{\tau_m (4Q^2 + 1)}{(s \tau_m - i \tau_m \omega_m) \left\{ (s \tau_c + 1)^2 + 4Q^2 \right\}}.$$  

(10)

The response $A(t)$ of the lock-in is the imaginary part of the inverse Laplace transform $L^{-1} \{a(s + i\omega_m)/(1 + \tau s)\}$ with $\gamma$ the lock-in time constant. An approximation can
be obtained in the special case where \( \omega_m = \omega_c \) in the limit \( Q \gg 1 \) and \( \omega_c \gg (1/\tau_m, 1/\tau_l) \).

\[
\frac{A(t)}{A_0} \approx \frac{1}{(1/t_m - 1/t_c)(\tau_c - \tau_l)} \exp \left(-\frac{t}{\tau_c}\right) + \frac{t_m}{\tau_m} \exp \left(-\frac{t}{\tau_l}\right) + \frac{t_m}{\tau_m} \exp \left(-\frac{t}{\tau_m}\right). \tag{11}
\]

The sum of these three exponentials vanishes at \( t = 0 \) and each term decays to zero with a different time constant at later time \( t > 0 \). This leads to a peaked lock-in signal \( A_{\text{peak}} \) whose amplitude and position depends on \( \tau_m \) (for a fixed \( \tau_c \) and \( \tau_l \)). At the end of the c.w. sequence in Fig.10 (\( t = 6s \)), the free oscillations decay of the cantilever (time constant \( \tau_l \)) are observed. If one tries to fit the data with the above nonlinear form, \( \tau_m = 2.2 \pm 0.07s \) is obtained but the quality of the fit is not very good. Values of \( \tau_m \) smaller than \( T_{1x} = 3.2s \) have also been reported by Verhagen et al. [21] and these findings were attributed to the phase noise of the r.f. source. However, when large modulation amplitude are employed for the c.w. sequence, such a fast force decay can also be consistent with a magnetization decrease due to a lack of adiabaticity.

To understand further the meaning of this fit parameter \( \tau_m \), we plot in Fig.11a the lock-in peak amplitude measured for different values of \( \Omega \) and \( B_1 \) when \( B_{\text{ext}} = 0.9425T \). The value of the non-adiabaticity parameter \( \alpha \) increases along the abscissa axis. For a fixed \( \Omega \) and \( B_1 \), the value of \( \alpha \) oscillates with time and passes through a maximum, \( \alpha_{\text{max}} = \Omega \omega_m / \gamma^2 B_1^2 \), at time \( t = 0 \) modulo \( \pi / \omega_m \). Fig.11b shows the amplitude of the peak signal \( A_{\text{peak}} \) predicted by equation (11) with \( F(t) \) calculated from equation (7) using a sample thickness of \( 7\mu m \). The results are normalized by \( A_{\text{total}} = Qg_M V_z / (kv_f^2) \) the amplitude associated with a uniform inversion of all spin inside the sample. The parameters introduced in the model are the values of the spin-lattice relaxation times measured on powder samples by conventional NMR with \( T_{1x} = 4.9s \) along the static field and \( T_{1x} = 3.2s \) along a 10G r.f. field. In the approximation that \( T_{1y} = T_{1x} \). In our theoretical model, perturbation effects from the dipolar broadening are also introduced. They are approximated as a time independent local field of Lorentzian lineshape. The solid lines are the shape calculated with \( T_2 = 40\mu s \) (the dotted lines correspond to \( T_2 = 100\mu s \)).

The increase of the force signal at small \( \Omega (\leq 50kHz) \) corresponds to an increase of the modulated magnetic moment. A larger frequency deviation increases the width of the probed slice, \( \Gamma \), and more protons oscillate at the frequency \( \omega_m \). For both large \( B_1 \geq 14G \) and large \( \Omega \geq 150kHz \) the amplitude of the signal eventually saturates when \( \Gamma \) becomes larger than the sample thickness.

For \( B_1 = 7 \) and 10G, the deviation from the low \( \Omega \)-linear increase (indicated by the arrows) marks the crossover from an adiabatic regime to a quasi-adiabatic one [40]. In our sample the threshold occurs at \( \alpha_{\text{max}} = 0.1 \), in good agreement with the theoretical model. In the adiabatic regime \( (\alpha_{\text{max}} < 0.1) \) the value of \( \tau_m \) is determined by \( T_{1\rho} \) effects, while in the quasi-adiabatic regime \( (0.1 < \alpha_{\text{max}} < 1) \) the magnetization decay \( \tau_m \) becomes smaller than \( T_{1x} \). The predicted position of this cross-over depends somewhat on the shape of the proton linewidth (dotted lines).

From the last discussion, one can conclude that the settings of Fig.10 correspond to a non-adiabatic parameter \( \alpha_{\text{max}} = 0.04 \) well inside the adiabatic regime for our compound. It can then be inferred that the decrease of the force signal in Fig.10 is due to spin-lattice effects in the rotating frame and our fitted value of \( \tau_m \) must be an average of the two \( T_1 \) reported in Fig.7.

The time dependence of the lock-in signal is fitted with a double damped synchronous excitation of two spin populations with respectively short \( (\tau_{mS}) \) and long \( (\tau_{ml}) \) relaxation times. The nonlinear function \( n_s A(t, \tau_{ms}) + (1 - n_s) A(t, \tau_{ml}) \) is used where \( A(t) \) is given by equation (11). The best fit is obtained for \( \tau_{ms} = 0.55 \pm 0.02s \), \( \tau_{ml} = 4.8 \pm 0.2s \) and \( n_s = 69 \pm 1% \). The result is the solid line shown in Fig.12. The fit values obtained for the relaxation times are similar to those measured in the magnetization recovery experiment in Fig.7. On Fig.12 the separate contribution to the lock-in signal of each spin species (dashed lines) are shown. The maximum force signal of the two spin species occurs respectively 0.7s and 1.9s after the start of the c.w. sequence for the short and long

![Fig. 11. (a) Measurements of the amplitude of the lock-in signal as a function of the modulation amplitude for different strength of the r.f. field. The dashed lines are guides for the eye. The arrows indicate the limit of the adiabatic regime. The measurements are performed at \( B_{ext} = 0.9425T \). (b) Calculated amplitude of the lock-in signal obtained for a 7\( \mu m \) thick sample using equation (7). The parameters introduced in the model are \( T_{1x} = 4.9s \), \( T_{1x} = 3.2s \) and \( T_2 = 40\mu s \) (solid line) or 100\( \mu s \) (dotted line).](image-url)
It brings up the question on how to define the lock-in peak amplitude in the case of a sample containing two spin species. We recall that our definition of the force signal is the average of the lock-in peak amplitude over a 1s time interval around its maximum value. This approach gives approximately equal weights to both spin species in the measurement. One can also observe in Fig.12 that the height of the two peaks are approximately equal despite the fact that there is 2.3 times more spin with short relaxation. As a matter of fact it can be shown that the mechanical detection is 2.4 times more sensitive to spin that have a 4.8s relaxation time compared to spin that have a 0.55s one. From this result, the value of the fit parameter \( n_s = 2.4 g_s / (2.4 - 1) g_s + 1 \) = 70\%, a value that agrees well with the fit \( n_a \) in Fig.12.

Finally, the expected amplitude of the force signal for our sample is calculated. Using equation (4), one gets a value of \( A_0 = 100 \text{Å} \) for the settings of the c.w. sequence used in Fig.12 (\( B_1 = 10 \text{G} \) and \( \Omega/2\pi = 50 \text{kHz} \)). In Fig.12 the predicted maximum of the lock-in peak is 0.3\( A_0 \), which is close to the experimentally measured value of 20\( \text{Å} \). In conclusion, our measured amplitude of the peak lock-in signal is in good agreement with the theoretical prediction if the two spin-lattice relaxation times of the two spin species are taken into account. Other effects such as misalignment of the sample compared to the cylinder magnetic axis can account partially for a decrease of the signal (e.g. a small offset of 0.1mm from the axis decreases the amplitude of the lock-in signal by a factor of 2).

5 Conclusion

Measurement sequences combining fast adiabatic passages and pulses are reported. They allow us to measure \( T_1 \) and \( T_2 \) for microscopic samples using a mechanical detection. This has been applied to quantitative analyses of the detected signals for a 7\( \mu \)m thick sample of \((\text{NH}_4)_2\text{SO}_4\). The transverse relaxation \( T_2 \) has been found consistent with conventional NMR detection on a macroscopic sample. Our sample displays however two spin lattice relaxation times \( T_1s = 0.4s \) and \( T_1l = 5s \). While the long \( T_1l \) corresponds to that measured for coarse powder samples, the short \( T_1s \) might be due to water contamination of our 7\( \mu \)m thick crystal during its contact with air. This contamination is found to be uniform in the thickness of the sample. This large difference in \( T_1 \) values has allowed us to study the influence of the spin-lattice relaxation in the rotating frame on the measured time dependence of the lock-in signal, as well as the variation of signal intensity with increasing non-adiabaticity of the sweep sequence. A consistent analysis of all the experimental parameters has been proposed and will be quite useful in future quantitative investigations of MRFM signals. Our work also raises the problem on how to perform reliable spin lattice relaxation measurements at different locations in the sample. Our investigation is mainly restrained to the bulk, i.e. the middle of the sample. It is known that the time dependence of the lock-in signal (and thus the apparent relaxation times) depends strongly on the value of \( B_{\text{ext}} \). These difficulties prevented us from interpreting quantitatively our results on the spatial distribution of the different spin densities close to the sample surface. This issue will be best resolved by performing a similar experiment on a hetero-layer sample of well-characterized composition.

We are greatly indebted to A. Dooglav for his help in the conventional NMR experiments. We also would like to thank C. Fermon, M. Goldman, J.F. Jacquinot and G. Lampel for stimulating discussions. This research was partly supported by the Ultimatech Program of the CNRS.

A Inhomogeneous field

Near the axis, a uniformly magnetized \((M_s)\) cylinder of length \( l \) and diameter \( \phi \) produces a field, whose component along \( k \), \( B_{\text{cyl}} \), decays radially as

\[
\frac{B_{\text{cyl}}(r, z)}{4\pi M_s} = \begin{pmatrix} b_{\frac{1}{2}} \left( \frac{z + l}{\phi} \right) - b_{\frac{3}{2}} \left( \frac{z}{\phi} \right) \\ + 3 \left( b_{\frac{1}{2}} \left( \frac{z + l}{\phi} \right) - b_{\frac{3}{2}} \left( \frac{z}{\phi} \right) \right) \frac{r^2}{\phi^2} + \mathcal{O}(r^4) \end{pmatrix}
\]

with \( b_{\frac{1}{2}}(z) = z(1 + 4z^2)^{-\frac{1}{2}} \). The fields are expressed in cylindrical coordinates with the origin centered on the cylinder upper surface (see Fig.1). In our case \( M_s \) is calculated from the applied field \( B_{\text{ext}} \) needed to produce a resonance signal at the sample position, \( z = 0.70 \text{mm} \). \( B_{\text{cyl}}(0, z = 0.70) = \omega_0 / \gamma - B_{\text{ext}} = 0.352 \text{T} \) is replaced in the above expression and one obtains \( M_s \approx 1400 \text{emu/cm}^3 \) for our iron. Using this result, the gradient \( g = -470 \text{T/m} \) at the sample location can be calculated.

![Fig. 12. Signal of Fig.10: The solid line is the response of an harmonic oscillator driven by two synchronous forces with short and long relaxations times \( n_s A(t, \tau_{ns}) + (1 - n_s) A(t, \tau_{nl}) \). The best fit is obtained for \( \tau_{ns} = 0.55 \pm 0.02s, \tau_{nl} = 4.8 \pm 0.2s \) and \( n_s = 70\% \). A value that agrees well with the fit used in Fig.12 (\( B \) value of our sample is calculated. Using equation (4), one gets a relaxation time compared to spin that have a 0.55s one. From this result, the value of the fit parameter \( n_s \) is 2.4\% \); a value that agrees well with the fit \( n_a \) in Fig.12. Finally, the expected amplitude of the force signal for our sample is calculated. Using equation (4), one gets a value of \( A_0 = 100 \text{Å} \) for the settings of the c.w. sequence used in Fig.12 (\( B_1 = 10 \text{G} \) and \( \Omega/2\pi = 50 \text{kHz} \)). In Fig.12 the predicted maximum of the lock-in peak is 0.3\( A_0 \), which is close to the experimentally measured value of 20\( \text{Å} \). In conclusion, our measured amplitude of the peak lock-in signal is in good agreement with the theoretical prediction if the two spin-lattice relaxation times of the two spin species are taken into account. Other effects such as misalignment of the sample compared to the cylinder magnetic axis can account partially for a decrease of the signal (e.g. a small offset of 0.1mm from the axis decreases the amplitude of the lock-in signal by a factor of 2). 5 Conclusion Measurement sequences combining fast adiabatic passages and pulses are reported. They allow us to measure \( T_1 \) and \( T_2 \) for microscopic samples using a mechanical detection. This has been applied to quantitative analyses of the detected signals for a 7\( \mu \)m thick sample of \((\text{NH}_4)_2\text{SO}_4\). The transverse relaxation \( T_2 \) has been found consistent with conventional NMR detection on a macroscopic sample. Our sample displays however two spin lattice relaxation times \( T_1s = 0.4s \) and \( T_1l = 5s \). While the long \( T_1l \) corresponds to that measured for coarse powder samples, the short \( T_1s \) might be due to water contamination of our 7\( \mu \)m thick crystal during its contact with air. This contamination is found to be uniform in the thickness of the sample. This large difference in \( T_1 \) values has allowed us to study the influence of the spin-lattice relaxation in the rotating frame on the measured time dependence of the lock-in signal, as well as the variation of signal intensity with increasing non-adiabaticity of the sweep sequence. A consistent analysis of all the experimental parameters has been proposed and will be quite useful in future quantitative investigations of MRFM signals. Our work also raises the problem on how to perform reliable spin lattice relaxation measurements at different locations in the sample. Our investigation is mainly restrained to the bulk, i.e. the middle of the sample. It is known that the time dependence of the lock-in signal (and thus the apparent relaxation times) depends strongly on the value of \( B_{\text{ext}} \). These difficulties prevented us from interpreting quantitatively our results on the spatial distribution of the different spin densities close to the sample surface. This issue will be best resolved by performing a similar experiment on a hetero-layer sample of well-characterized composition. We are greatly indebted to A. Dooglav for his help in the conventional NMR experiments. We also would like to thank C. Fermon, M. Goldman, J.F. Jacquinot and G. Lampel for stimulating discussions. This research was partly supported by the Ultimatech Program of the CNRS. A Inhomogeneous field Near the axis, a uniformly magnetized \((M_s)\) cylinder of length \( l \) and diameter \( \phi \) produces a field, whose component along \( k \), \( B_{\text{cyl}} \), decays radially as

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\]

with \( b_{\frac{1}{2}}(z) = z(1 + 4z^2)^{-\frac{1}{2}} \). The fields are expressed in cylindrical coordinates with the origin centered on the cylinder upper surface (see Fig.1). In our case \( M_s \) is calculated from the applied field \( B_{\text{ext}} \) needed to produce a resonance signal at the sample position, \( z = 0.70 \text{mm} \). \( B_{\text{cyl}}(0, z = 0.70) = \omega_0 / \gamma - B_{\text{ext}} = 0.352 \text{T} \) is replaced in the above expression and one obtains \( M_s \approx 1400 \text{emu/cm}^3 \) for our iron. Using this result, the gradient \( g = -470 \text{T/m} \) at the sample location can be calculated.
B Adiabaticity

The aim of this appendix B is to calculate the decrease of the magnetization due to the spin-lattice relaxation and the lack of adiabaticity. The solution below is proposed by M. Goldman. In the limit of strong r.f. fields (larger than the local field), one can neglect the spin-lattice relaxation of the dipolar energy expectation value. In the rotating frame, the time evolution of the different spin components are [40]:

\[
\frac{\partial \langle I_z \rangle}{\partial t} = -\gamma B_1 \langle I_y \rangle + \frac{\langle I_0 \rangle - \langle I_z \rangle}{T_{1z}} \tag{B.1a}
\]

\[
\frac{\partial \langle I_x \rangle}{\partial t} = +\gamma B_1 \cot \theta \langle I_y \rangle - \frac{\langle I_y \rangle - \langle I_z \rangle - \langle I_x \rangle}{\tau_s} \tag{B.1b}
\]

\[
\frac{\partial \langle I_y \rangle}{\partial t} = +\gamma B_1 ((\langle I_z \rangle - \cot \theta \langle I_x \rangle) - \frac{\langle I_y \rangle - \langle I_z \rangle - \langle I_x \rangle}{\tau_s} - \frac{i}{\hbar} \langle [H_{DZ}, I_y] \rangle \tag{B.1c}
\]

with \( \langle I \rangle = \text{Tr}(I\sigma) \) the expectation value of the magnetization, \( \sigma \) the instantaneous density matrix in the rotating frame and \( H_{DZ} \) the secular part of the dipolar Hamiltonian. The commutator incorporates the local field contribution defined through \( B^2 = \text{Tr}(H_{DZ}^2)/\gamma^2 \text{Tr}(I_z^2) \). In our notation \( \theta \) is the angle between the directions of the static and effective field, \( B_e = B_1/\sin \theta \), with \( B_1 \cot \theta \) the projection along \( \mathbf{k} \). Under r.f. irradiation, a new coordinate system \( \{X, Y, Z\} \) is defined through the unitary operator \( \exp(-i \theta I_y) \), a rotation around \( y \) by \( \theta \). In the doubly rotating frame the differential equations then become:

\[
\frac{\partial \langle I_z \rangle}{\partial t} = +\gamma \frac{B_1}{T_{1z}} \frac{s^2}{T_{1z}^2} \left( \frac{\langle I_0 \rangle - \langle I_z \rangle}{T_{1z}} \right) \left( \langle I_z \rangle - \cot \theta \langle I_x \rangle \right) - \frac{i}{\hbar} \langle [H_{DZ}, I_y] \rangle \tag{B.2a}
\]

\[
\frac{\partial \langle I_x \rangle}{\partial t} = -\gamma \frac{B_1}{T_{1z}} \left( \frac{\langle I_0 \rangle - \langle I_x \rangle}{T_{1z}} \right) - \frac{i}{\hbar} \langle [H_{DZ}, I_x] \rangle \tag{B.2b}
\]

\[
\frac{\partial \langle I_y \rangle}{\partial t} = -\gamma \frac{B_1}{T_{1z}} \left( \frac{\langle I_0 \rangle - \langle I_y \rangle}{T_{1z}} - \frac{3c^2 - 1}{2} \right) \left( \langle I_z \rangle - \cot \theta \langle I_x \rangle \right) - \frac{i}{\hbar} \langle [H_{DZ}, I_y] \rangle \tag{B.2c}
\]

with \( s = \sin \theta \) and \( c = \cos \theta \). \( H_{DZ} \) is the doubly truncated part of the dipolar Hamiltonian that commutes with \( I_z \). The term \( \dot{\theta} \gamma \dot{\theta} \) is the inertia field due to the transformation to a time-dependent reference axis. In the adiabatic regime, defined by \( \alpha = \dot{\theta} / (\gamma B_e) \ll 1 \), it can be shown that \( \langle I_x \rangle = \langle I_y \rangle = 0 \) and the first two terms of equation (B.2a) are the expression of the spin-lattice relaxation in the rotating frame for strong r.f. fields \([1,41]\). This appendix seeks to evaluate the term proportional to \( \langle I_x \rangle \) in equation (B.2a) that represents the decrease of \( \langle I_z \rangle \) due to the lack of adiabaticity. Our investigation will be restricted to the quasi-adiabatic regime \([40]\), where non-adiabaticity corrections might be dominant over \( T_{1p} \) effects but \( \alpha < 1 \). The equations of motion are expressed in terms of the raising and lowering operators \( I^+ = (I_X + iI_Y) \) and \( I^- \) its complex conjugate. We suppose that \( \gamma B_e \ll 1/\tau_s \), the reorientation correlation rate, for our compound which leads to \( T_{1z} \) independent of \( \theta \) and \( T_{1y} \approx T_{1z} \). As a further simplification, spin-spin interactions are neglected and these effects will be calculated numerically later on. As a consequence, one has

\[
\frac{\partial \langle I^+ \rangle}{\partial t} \approx -i\gamma \frac{B_1}{s} \langle I_0 \rangle - \langle I^+ \rangle - \frac{\dot{\theta}}{T_{1z}} - \frac{\dot{\theta} \langle I_0 \rangle}{T_{1z}} - \frac{1}{T_{1z}} \left( \frac{1}{T_{1z}} \right) - \frac{1}{T_{1z}} \langle I_Z \rangle \tag{B.3}
\]

with \( 1/T_{1z}^2 = (1/T_{1x} + c^2/T_{1z} + s^2/T_{1z})/2 \). Furthermore, the period of the cyclic passage is much smaller than the spin-lattice relaxation times. Hence, both \( 1/T_{1z} \) and \( 1/T_{1x} \) are negligible compared to \( \dot{\theta} \):

\[
\frac{\partial \langle I^+ \rangle}{\partial t} \approx -i\gamma \frac{B_1}{\sin \theta} \langle I_0 \rangle - \langle I^+ \rangle - \dot{\theta} \langle I_0 \rangle - \sin \theta \langle I_0 \rangle \tag{B.4}
\]

which, upon integration, gives the result:

\[
\langle I^+ \rangle = -\int_0^t \left[ \dot{\theta}(t') \langle I_0 \rangle + \sin \theta(t') \langle I_0 \rangle \right] \exp \left( -\int_t^t \frac{1}{T_{1z}} \frac{1}{T_{1z}} \frac{1}{T_{1z}} \right) \frac{1}{T_{1z}} \langle I_0 \rangle \langle I_0 \rangle \tag{B.5}
\]

assuming that \( \langle I^+ \rangle = 0 \) at a time \( t = 0 \). The expression for \( \langle I^- \rangle \) is the complex conjugate of the above expression. For values of \( t < T_1 \), it is a good approximation to neglect \( \langle I_0 \rangle/T_{1z} \) compared to \( \dot{\theta} \langle I_0 \rangle \) in the first bracket. Since the decay of \( \langle I_Z \rangle \) is slow, it is replaced by a constant. Finally, the time variation of the longitudinal magnetization is given by:

\[
\frac{\partial \langle I_z \rangle}{\partial t} \approx -\langle I_z \rangle \dot{\theta}(t) \int_0^t \dot{\theta}(t') \exp \left( -\int_t^t \frac{1}{T_{1z}} \frac{1}{T_{1z}} \right) \frac{1}{T_{1z}} \frac{1}{T_{1z}} \frac{1}{T_{1z}} \right) \tag{B.6}
\]

which has been used in equation (6) in the text. Although the general form of our final expression contains a \( T_{1z}^{-1} \) exponential decay in the first integral, the decrease of the magnetization due to the quasi-adiabatic part of the c.w. sequence is unrelated to spin-lattice relaxation mechanisms. This is best seen in the free spin limit \( T_1 \rightarrow \infty \), where the first term in equation (B.6) reduces to the integral form of the ordinary differential equation \( \partial \langle I \rangle/\partial t = \gamma \langle I \rangle \times B_e \), an equation of motion that preserves the norm of the magnetization. Our above solution expresses the
The new system of differential equations is obtained by applying the unitary transformation $\exp(-i\alpha I_X)$ to the equation system (B.2). Here second order corrections are included, the amplitude $\langle I_Z \rangle$ decays toward zero and the expectation value of the norm of the magnetization follows approximately the instantaneous value of $\langle I_Z \rangle$.

References

1. A. Abragam, *Principles of Nuclear Magnetism* (Oxford University Press, 1961).
2. J. A. Sidles, Appl. Phys. Lett. 58, 2854 (1991).
3. J. A. Sidles, Phys. Rev. Lett. 68, 1124 (1992).
4. J. A. Sidles and D. Rugar, Phys. Rev. Lett. 70, 3506 (1993).
5. J. A. Sidles, J. L. Garbini, K. J. Bruland, D. Rugar, O. Züger, S. Hoen, and C. S. Yannoni, Rev. Mod. Phys. 67, 249 (1995).
6. D. Rugar, C. S. Yannoni, and J. A. Sidles, Nature 360, 563 (1992).
7. D. Rugar, O. Züger, S. Hoen, C. S. Yannoni, H. M. Vieth, and R. D. Kendrick, Science 264, 1560 (1994).
8. Z. Zhang, P. C. Hammel, and P. E. Wigen, Appl. Phys. Lett. 68, 2005 (1996).
9. O. Züger and D. Rugar, Appl. Phys. Lett. 63, 2496 (1993).
10. P. C. Hammel, Z. Zhang, G. J. Moore, and M. L. Roukes, J. Low Temp. Phys. 101, 59 (1995).
11. O. Züger, S. T. Hoen, C. S. Yannoni, and D. Rugar, J. Appl. Phys. 79, 1881 (1996).
12. A. Schaff and W. S. Veeman, Appl. Phys. Lett. 70, 2598 (1997).
13. B. J. Suh, P. C. Hammel, Z. Zhang, and M. M. M. et al., J. Vac. Sci. and Technol. B 16, 2275 (1998).
14. C. Ascoli, P. Baschieri, C. Frediani, L. Lenci, M. Martinelli, G. Alzetta, R. M. Celi, and L. Pardi, Appl. Phys. Lett. 69, 3020 (1996).
15. K. Wago, O. Züger, R. Kendrick, C. S. Yannoni, and D. Rugar, J. Vac. Sci. and Technol. B 14, 1197 (1996).
16. K. Wago, O. Züger, J. Weneger, R. Kendrick, C. S. Yannoni, and D. Rugar, Rev. Sci. Instrum. 68, 1823 (1997).
17. K. Wago, D. Botkin, C. S. Yannoni, and D. Rugar, Phys. Rev. B 57, 1108 (1998).
18. K. Wago, D. Botkin, C. S. Yannoni, and D. Rugar, Appl. Phys. Lett. 72, 2757 (1998).
19. K. J. Bruland, W. M. Dougherty, J. L. Garbini, J. A. Sidles, and S. H. Chao, J. Appl. Phys. 73, 3159 (1998).
20. A. Schaff and W. S. Veeman, J. Magn. Res. 126, 200 (1997).
21. R. Verhagen, C. W. Hilbers, A. P. M. Kentgens, L. Lenci, R. Groeneveld, A. Wittli, and H. van Kempen, Phys. Chem. Chem. Phys. 1, 4025 (1999).
22. S. R. Miller, R. Blinc, M. Brennan, and J. S. Waugh, Phys. Rev. 126, 528 (1962).
23. D. E. O’Reilly and T. Tsang, J. Chem. Phys. 46, 1291 (1967).
24. S. Guillous and O. Klein (to be published).
25. Near the symmetry axis, $\nabla B_{rad} \approx \partial B_{rad}/\partial z$.
26. 99.99% purity (Aldrich).
27. The value is measured in a 1T field using a coil detection sensor, we estimate to +1◦C the elevation of the sample temperature during the r.f.-irradiation.
28. R. E. Richards and T. Scaeer, Trans. Faraday Soc. 57, 210 (1961).
29. The sample mass and shape are stable for the duration of the experiment.
30. Using the static flexion of the cantilever as a temperature sensor, we estimate to +1.5°C the elevation of the sample temperature during the r.f.-irradiation.
31. We can do this approximation because all slices are phase coherent.
32. The width of the probed slice is proportional to $\sqrt{1.2/(\gamma b)^2 + B_1^2/g}$.
33. The fast relaxation $T_{1s}$ can also partially explain the positive offset $b$.
34. In the limit $B_1 > g \times$thickness, $\tau^{-1}$ equals to $\left(1/T_1 + 1/2T_2\right)/2$ [1].
35. E. Fukushima and S. B. W. Roeder, *Experimental pulse NMR* (Addison-Wesley, 1981).
36. E. O. Schlemper and W. C. Hamilton, *J. Chem. Phys.* 44, 4498 (1966).
37. J. L. Hutter and J. Bechhoefer, Rev. Sci. Instrum. 64, 1868 (1993).
38. Comments about MRFM noise can be found in http://weber.u.washington.edu/~sidles/.
39. The phase noise of our setup is estimated from fast adiabatic passage measurements on de-ionized H2O in a 1T field with a coil detection and comparing the apparent $\tau_m$ obtained with the VCO and a Marconi 2023 synthesizer.
40. M. Goldman, *Spin temperature and Nuclear Magnetic Resonance in solids* (Oxford University Press, 1970).
41. C. P. Slichter, *Principles of Magnetic Resonance* (Springer, Berlin, Heidelberg, New York, 1996), 3rd ed.