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Long rotational coherence times of molecules in a magnetic trap

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Polar molecules in superpositions of rotational states exhibit long-range dipolar interactions, but maintaining their coherence in a trapped sample is a challenge. We present calculations that show many laser-coolable molecules have convenient rotational transitions that are exceptionally insensitive to magnetic fields. We verify this experimentally for CaF where we find a transition with sensitivity below 5 Hz G−1 and use it to demonstrate a rotational coherence time of 6.4(8) ms in a magnetic trap. Simulations suggest it is feasible to extend this to more than 1 s using a smaller cloud in a biased magnetic trap.

Ultracold polar molecules present exciting opportunities for quantum simulation, quantum computation, and tests of fundamental physics [1]. The rotational motion is particularly important for these applications. Rotational transitions are easily driven by microwave fields, and excited rotational states have long lifetimes. Polar molecules in superpositions of rotational states have large oscillating electric dipole moments, providing long-range dipole-dipole interactions between them [2–4]. The strong electric dipole coupling of rotational states to microwave fields can also be used to interface gas-phase molecules with mesoscopic solid-state systems [5, 6]. Many schemes have been proposed to use these interactions to generate entanglement, engineer inter-particle potentials and implement two-qubit quantum gates [7–11].

Most applications require trapped molecules and coherence times that are long compared to the characteristic interaction strength. Here, we explore how that can be achieved with molecules in magnetic traps. To avoid the dephasing that arises from an inhomogeneous transition frequency, the trap potential should, ideally, be identical for the two rotational states. Experiments so far have focused exclusively on optical traps in the form of optical lattices or arrays of tweezer traps where inter-particle separations below 1 μm are possible, producing dipole-dipole couplings with energies ~ 1 kHz. Whilst second-long coherence times have been demonstrated for superpositions of hyperfine states [12], rotational coherence times longer than ~ 1 ms are a challenge because of the dependence of the ac Stark shift on rotational state [13]. By using a dc electric field to uncouple the rotational angular momentum from the nuclear spin, and setting the polarization angle of the trapping light to equalize the Stark shifts of the chosen rotational states, Seefelberg et al. [14] extended rotational coherence times to 8.7(6) ms in a sample of optically trapped NaK molecules. Long vibrational coherence times have also been demonstrated for Sr2 molecules in a state-insensitive optical lattice [15].

Recently, direct laser cooling has produced ultracold molecules with both electric and magnetic moments [16–19], opening new possibilities for experiments in magnetic traps [20, 21]. Two-dimensional arrays of magnetic traps have been demonstrated for atoms with spacings comparable to those achievable with optical arrays [22]. Molecules might also be held in magnetic chip traps close to superconducting microwave resonators [5], where the regime of strong coupling between molecules and microwave photons may be reached, and where molecules may be coupled via the resonator with interaction strengths of order 100 kHz.

To obtain long rotational coherence times in a magnetic trap, it is necessary to find states with large and nearly identical magnetic moments. Here, we investigate rotational transitions in 2Σ molecules and find a pair of states in CaF with magnetic moments of ~ 1 μB that are equal to 3.3(1) parts per million. We demonstrate a superposition of the two states in a quadrupole magnetic trap with a coherence time of 6.4(8) ms. By comparison with simulations and free-space measurements we establish the dominant dephasing mechanisms and propose ways to achieve coherence times exceeding 1 s.

The interaction of a 2Σ molecule with a magnetic field $\vec{B}$ is described by the Zeeman Hamiltonian [23, 24],

$$H_Z = H_e + H_a + H_r + H_a = g_s \mu_B \vec{S} \cdot \vec{B} - \sum_i g_i \mu_N \vec{I}_i \cdot \vec{B} - g_e \mu_B \vec{N} \cdot \vec{B} + g_l \mu_B (\vec{S} \cdot \vec{B} - (\vec{S} \cdot \hat{z})(\vec{B} \cdot \hat{z})), \quad (1)$$

where $\hat{z}$ is a unit vector in the direction of the internuclear axis, $\vec{S}$ is the electron spin operator, $\vec{N}$ is the rotational angular momentum operator, $\vec{I}_i$ is the spin operator of nucleus $i$, and the sum is over the nuclei. $H_e$ describes the interaction of the electron magnetic moment with $\vec{B}$ and is, by far, the largest term. $H_a$ describes the much smaller contribution from the nuclear magnetic moments, $H_r$ the rotational Zeeman interaction, and $H_l$ the anisotropic correction to the electronic Zeeman interaction. The search for magnetically insensitive rotational transitions is hindered by the Hamiltonian for the fine and hyperfine structure, $H_{\text{hf}}$, which couples
the angular momenta in a way that depends on \(N\) [23]. This is solved by choosing the stretched states \(|N\rangle_{\text{str}} = |N, m_N = N\rangle |S, m_S = S\rangle |I, m_I = I\rangle\), where \(m_X\) is the projection of \(X\) onto the magnetic field axis. These states are eigenstates of \(H_{\text{Hfr}}\) and \(H_Z\). Their Zeeman shifts, \(\Delta E_N\), are almost identical, because the large contribution from \(H_e\), and the smaller one from \(H_n\), are both independent of \(N\). For the single-photon transitions \(|N\rangle_{\text{str}} \leftrightarrow |N + 1\rangle_{\text{str}}\), the residual magnetic sensitivity due to the two remaining terms is

\[
\Delta \mu(N) = \frac{(\Delta E_{N+1} - \Delta E_N)}{B} = \frac{g_I - g_L}{(2N + 4)^2} - g_r \mu_B. \tag{2}
\]

If the ratio \(g_I/g_r\) is close to \((2N + 4)^2 - 1\) for some \(N\), the remaining two terms nearly cancel, giving the desired magnetic insensitivity.

We have carried out calculations of this ratio for a variety of alkaline-earth fluorides, hydrides and hydroxides of interest for laser cooling. To evaluate \(\mu\) for transitions \(|N\rangle_{\text{str}} \leftrightarrow |N + 1\rangle_{\text{str}}\) up to \(N = 6\). For \(N = 0\), the sensitivities are dominated by \(g_r\), and for large \(N\) they approach the value set by \(g_I\). The hydrides generally have much larger values of both \(g_I\) and \(g_r\) and have no transitions with residual sensitivities on the scale shown. The remaining molecules each exhibit at least one transition with \(|\Delta \mu/h| < 30\,\text{Hz}\,\text{G}^{-1}\) and six of them have a transition with \(|\Delta \mu/h| < 10\,\text{Hz}\,\text{G}^{-1}\). These results suggest that convenient rotational transitions with extremely small magnetic sensitivities are a common feature of alkaline-earth fluorides and hydroxides.

We have used Ramsey spectroscopy in a uniform magnetic field \(B\) to determine the magnetic sensitivities of such transitions experimentally in \(\text{CaF}\). We measure \(\Delta \mu\) for two transitions, \(|0\rangle_{\text{str}} \leftrightarrow |1\rangle_{\text{str}}\) and \(|1\rangle_{\text{str}} \leftrightarrow |2\rangle_{\text{str}}\), whose frequencies are \(f_{0\leftrightarrow1} = 20\,553.4\,\text{MHz}\) and \(f_{1\leftrightarrow2} = 41\,088.9\,\text{MHz}\). We capture \(\sim 10^4\) molecules in a magneto-optical trap [17], cool them to about \(50\,\mu\text{K}\) [48], then prepare them in a single internal state, either \(|0\rangle_{\text{str}}\) or \(|1\rangle_{\text{str}}\) [21]. The Ramsey sequence [49] uses a pair of nearly resonant \(\pi/2\) pulses, \(20\,\mu\text{s}\) long, separated by a free-evolution time \(\tau\). Following the second pulse, we measure the number of molecules in \(N = 1\) by recapturing them into the MOT and imaging their fluorescence. Dividing by the number initially loaded in the MOT gives a signal that is insensitive to shot-to-shot fluctuations in the number of molecules. The magnetic field at the molecules is calibrated to better than 1%
by single-pulse microwave spectroscopy of the transition \( |N = 0, F = 1, m_F = -1 \rangle \leftrightarrow |N = 1, F = 0 \rangle \). The Zeeman shift of this transition can be accurately calculated because it is dominated by \( H_{\text{int}} \), and because \( g_S \) and the hyperfine parameters are known to high precision [50].

Figure 2(a) shows Ramsey fringes for the transition \( 1_{\text{str}} \leftrightarrow 2_{\text{str}} \), obtained by scanning the microwave frequency. The dark blue data are for \( \tau = 4 \) ms and the light blue for \( \tau = 5 \) ms. Repeating the measurement with different \( \tau \) identifies the fringe corresponding to the centre frequency. Fitting to the fringes determines the transition frequency with statistical uncertainty below 1 Hz.

Figure 2(b) shows the change in transition frequency for both transitions as a function of \( B \). The solid lines show linear fits to the data that give residual magnetic sensitivities \( \Delta \mu / \hbar = -194(4) \) Hz G\(^{-1}\) for the transition \( 0_{\text{str}} \leftrightarrow 1_{\text{str}} \), and \(-4.7(2) \) Hz G\(^{-1}\) for \( 1_{\text{str}} \leftrightarrow 2_{\text{str}} \). The uncertainties are dominated by drifts in \( B \) between measurements of the insensitive transition and the transition used for calibration, represented by the horizontal error bars. The error in magnetic field is correlated between different points, because the drift is slow compared to the measurement time; this has been taken into account in the calculation of the uncertainties in \( \Delta \mu \). From these measurements we determine \( g_l = -1.87(8) \times 10^{-3} \) and \( g_r = -5.0(2) \times 10^{-5} \), in excellent agreement with the calculated values. At \( N = 1 \) the two terms in Eq. 2 cancel to within 3%, resulting in cancellation of the magnetic moments of the two states to 3.3(1) parts per million.

The magnetic insensitivity of the transition \( 1_{\text{str}} \leftrightarrow 2_{\text{str}} \) suggests that long coherence times should be possible in a magnetic trap. To investigate this we measure the decay of the beat note between the oscillations of the rotational superposition and those of a slightly detuned microwave field. The first \( \pi/2 \) pulse is applied and then the quadrupole trap is immediately turned on with an axial field gradient of 45 G cm\(^{-1}\). The lifetime of the molecules in the magnetic trap is about 4.5 s [21], much longer than the timescale of any of the experiments reported here. The second \( \pi/2 \) pulse is applied in the trap and the molecules are then recaptured and imaged in a MOT, as before. Figure 3(a) shows the Ramsey fringes traced out by scanning the free evolution time \( \tau \). The solid line is a fit to a sinusoid with exponentially decaying amplitude, which gives a coherence time of 6.4(8) ms.

To elucidate the sources of decoherence in the trap, we use a Monte-Carlo simulation of the Ramsey experiment in the magnetic trap. The result of this simulation fits well to an exponentially decaying sinusoid with \( 1/e \) time of 6.1 ms, in agreement with our experimental data. We identify three decoherence mechanisms: (i) The residual magnetic sensitivity of the transition causes a spread in the phase accumulated by different molecules.

(ii) The movement of a molecule between the two \( \pi/2 \) pulses causes a change of phase which is different for each molecule; this is a form of Doppler broadening. (iii) The molecules adiabatically follow the local magnetic field vector as they move along different trajectories. This last mechanism has two effects. The first is to vary the amplitude of the microwave field in the relevant polarisation for each molecule at each Ramsey pulse, resulting in a loss of contrast, but no longer-term dephasing. The second is to impart a geometric phase to each molecule, which depends on its trajectory between the two pulses, and is proportional to the difference in \( m_F \) between the two states [51]. To quantify the contributions of each mechanism, we artificially remove the effect of the others from the simulation and find the time taken for the amplitude of the Ramsey fringes to decay to a factor of \( e \). We find the decoherence rates for each mechanism to be: (i) 60 s\(^{-1}\), (ii) 50 s\(^{-1}\) and (iii) 100 s\(^{-1}\).

The decoherence rate of mechanism (i) scales linearly with the average field experienced by the molecules in the trap, currently set almost entirely by the initial Gaussian width of 1.5 mm. In the limit of a small initial cloud, the sample explores a range of fields determined only by its temperature, \( T \). The resulting spread of transition frequencies, \((k_B T / \hbar)(\Delta \mu / \mu_0)\), gives a decoherence rate of 0.14 \( \mu \)K\(^{-1}\) s\(^{-1}\). This implies an achievable coherence time of 1.4 s at 5 \( \mu \)K, a temperature that has already been demonstrated for CaF [52, 53]. The effect of mechanism (ii) is also related to the cloud size. When the maximum distance that a molecule moves is less than the wavelength of the microwave field, the phase change due to motion is limited to less than \( 2\pi \) and the superposition never fully decoheres. The result is a fixed loss of contrast at late times. In our experiment, this maximum phase change is set by the initial cloud size; reducing the cloud size would decrease the size of the effect. Mechanism (iii), currently the largest contribution to the decohe-
molecule along the $k$ vector is constant so the phase accumulated changes linearly with time. Consequently, this decoherence can be reversed by employing a spin-echo sequence in which a $\pi$ pulse is applied mid-way between the two $\pi/2$ pulses. Figure 3(c) shows Ramsey fringes in free space with a $\pi$ pulse applied 14.7 ms after the first $\pi/2$ pulse. Molecules at different positions experience different microwave powers, so a $\pi$ or $\pi/2$ pulse cannot be perfect for all molecules. The extra $\pi$ pulse for the spin-echo technique thus causes an overall reduction in contrast. The signal also falls as the molecules expand and drop out of the detection region. However, we see no decoherence at all on this timescale; our data restrict the magnitude of any effect to less than 20 s$^{-1}$ at the 95% confidence level.

The exceptionally small magnetic sensitivity of the transition $|1\rangle_{str} \leftrightarrow |2\rangle_{str}$ raises the question of whether the sensitivity might be even smaller in an excited vibrational state, $v > 0$. To consider this, we expand the calculated $g_l$ and $g_r$ for CaF as a Taylor series in the bond length $R$ around the equilibrium value $R_e$. For a harmonic oscillator, only even-order terms in this expansion contribute to expectation values. For a slightly anharmonic oscillator such as most chemically bound molecules, both the linear and quadratic terms contribute significantly to the dependence on $v$. We calculate $g_l$ and $g_r$ at two additional bond lengths, close to the inner and outer turning points for the $v = 1$ state of CaF, to evaluate the coefficients linear and quadratic in $(R - R_e)$. Combining these with the expectation values of $(R - R_e)$ and $(R - R_e)^2$ for CaF yields

$$
g_l(v) = -1.86 \times 10^{-3} \left[1 + 6.20 \times 10^{-3} \left(v + \frac{1}{2}\right) + \ldots\right];$$

$$
g_r(v) = -5.13 \times 10^{-5} \left[1 - 5.77 \times 10^{-3} \left(v + \frac{1}{2}\right) + \ldots\right];$$

$$
g_l(v)/g_r(v) = 36.16 \left[1 + 0.0120 \left(v + \frac{1}{2}\right) + \ldots\right]. \quad (3)$$

This indicates that the dependence on $v$ is small, and that the magnetic sensitivity is lowest for $v = 0$. Nevertheless, the magnitude of the effect suggests that vibrational state dependence may be useful for fine-tuning magnetic sensitivity in other molecules.

In conclusion, we have shown that many laser-coolable molecules have pairs of states belonging to neighbouring rotational manifolds whose magnetic moments are large and identical to a few parts per million. In CaF, the transition between the stretched states of $N = 1$ and $N = 2$ has a magnetic sensitivity of $-4.7(2)$ Hz G$^{-1}$. We have demonstrated a coherence time of 6.4(8) ms for a magnetically trapped sample in a superposition of these states. Characterisation of the principal sources of decoherence suggests that modifications to the trapping geometry and reduction of the cloud size will yield considerable improvements, with the prospect of coherence times greater than 1 s for small clouds at 5 µK or below. These results open up potential new platforms for quantum simulation and computation with polar molecules in magnetic microtraps [22] and provide a route to the regime of strong coupling between solid-state systems and molecules held...
in magnetic chip traps [5]. These magnetically insensitive transitions may also be valuable in precision measurement [55] where sensitivity to magnetic fields is an important source of systematic error.

Underlying data may be accessed from Zenodo [56] and used under the Creative Commons CCZero license. We thank J. Dyne and V. Gerulis for expert technical assistance. This work was supported by EPSRC under grants EP/M027716/1 and EP/P01058X/1, and by the Royal Society.

L. Caldwell and H.J. Williams contributed equally to this work.

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Long rotational coherence times of molecules in a magnetic trap:
Supplemental Material

I. ELECTRONIC STRUCTURE CALCULATIONS OF $g_r$ AND $g_l$

We calculate $g_l$, the anisotropy of the electronic $g$ tensor, using density functional theory (DFT) with the B3LYP functional [26]. This functional was chosen because, as discussed in reference [33], it is better than other functionals at reproducing experimental values of $g_l$ and $\gamma$; the latter characterizes the electron spin-rotation interaction and is connected to $g_l$ via Curl’s approximation [34]. The calculations include relativistic corrections via the zero-order regular approximation (ZORA) [28–30] and are performed with the ORCA code [25]. Experimental values of $g_l$ are available for numerous $^2\Sigma$ molecules, including most of those discussed in Table I [35–44, 50]. In general, our calculations agree with experimental values within 5%, although the difference may increase up to 20% when comparing with measurements performed in noble-gas matrices or when the value of $g_l$ is not directly measured but is extracted from $\gamma$ using Curl’s approximation, which is known to introduce an error around 15% [45]. Such larger discrepancies occur, for example, for MgOH (measured in a matrix [42]) and CaH ($g_l$ obtained from $\gamma$ [39]).

We are unable to use the same methodology for the rotational $g$ factor, $g_r$, because its calculation is not supported by ORCA. As correlation effects are relatively small for $g_r$ [46], we calculate it using Hartree-Fock (HF) calculations performed with the DALTON code [31]. Unfortunately, very few experimental values of $g_r$ are available for open-shell molecules and, in particular, for those included in Table I. The only exception to this is CaF, for which the calculated value is in good agreement with the experimental value obtained in the present work, $g_r = -5.0(2) \times 10^{-5}$.

We initially attempted to perform calculations on Yb as well as the lighter metals in Table I, but ultimately decided they were not reliable enough for publication. To provide consistency across all the elements, we chose to use the x2c-TZVPP basis sets of Pollak and Weigend [27] for both DFT and HF calculations. These basis sets are of triple-$\zeta$ quality, and are designed for use in relativistic calculations. They are available for all atoms in the periodic table up to Rn.

The DFT and HF methods we use here are reliable only for molecules that are reasonably well described by a single electron configuration. In order to verify this, we performed CCSD(T) calculations and used the T1 diagnostic [47] to assess the multiconfigurational character of the ground electronic state. A value $T1 > 0.02$ is generally considered to indicate that multireference calculations are needed. For the molecules in Table I, T1 was below this threshold, indicating that the ground state is reasonably well described by a single configuration.