Optical Excitation of Nuclear Spin Coherence in Tm$^{3+}$:YAG

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A thulium-doped crystal is experimentally shown to be an excellent candidate for broadband quantum storage in a solid-state medium. For the first time, nuclear spin coherence is optically excited, detected and characterized in such a crystal. The lifetime of the spin coherence – the potential storage entity – is measured by means of Raman echo to be about 300 µs over a wide range of ground state splittings. This flexibility, attractive for broadband operation, and well fitted to existing quantum sources, results from the simple hyperfine structure, contrasting with Pr- and Eu-doped crystals.

So far, the mapping of a quantum state of light onto an atomic ensemble has been implemented in atomic vapors [1] and cold atom clouds [2]. Because of their long optical coherence lifetimes at low temperature, rare-earth ion-doped crystals (REIC) have been extensively investigated for optical data storage [3] and data processing [4,5]. There has recently been renewed interest in these materials, stimulated by proposals to explore their adequacy to quantum memories [6, 7, 8, 9, 10]. To some extent, REIC at low impurity concentration are similar to atomic vapors with the advantage of no atomic diffusion.

Most optical quantum memory protocols rely on the transfer of a quantum state of light into a long-lived atomic spin coherence that is free from decoherence via spontaneous emission. This can be achieved in a Λ-type three-level system where two hyperfine or spin sublevels are optically connected to a common upper level. The presence of a Λ-system ensures efficient coupling between light and matter together with long storage times. A Λ-system also represents a basic device where, in a simple way, the transition to be excited by the quantum field can be triggered for storage or restitution by an external control field. In the prospect of quantum storage, electromagnetically induced transparency (EIT) protocols involving Λ-type have been studied in praseodymium-doped materials [6,7].

It is noteworthy that the essence of a Λ-system operation, namely the optical excitation of nuclear spin coherence, has been practised in REIC for almost 30 years [11,12,13,14,15,16], but always in praseodymium- or europium-doped crystals. However, quantum storage demonstration in Pr- or Eu-doped compounds is limited by the smallness of their hyperfine structure, not really matching the bandwidth of existing quantum sources. We recently demonstrated the existence of a Λ-system with adjustable ground state splitting in thulium-doped YAG [17]. Widely adjustable splitting might help to match the bandwidth of existing quantum sources. In this Letter, for the first time to the best of our knowledge, we optically excite, characterize and detect the nuclear spin coherence in the electronic ground state of a Tm-doped crystal.

In crystals doped with Pr$^{3+}$ or Eu$^{3+}$, Λ-systems are built on the hyperfine structure of the ground level with a sublevel splitting up to a few tens of MHz. This spacing represents the memory bandwidth. Indeed, the two transitions of the Λ cannot be polarization selected and thus only differ by their frequency. Therefore, in order to excite only the relevant single transition, the incoming signal must be spectrally narrower than the splitting. Applying an external magnetic field would increase the splitting and the memory bandwidth but would split each hyperfine level into two nuclear spin sublevels, thus drastically complexifying the level system. Besides, only dye lasers are available at Pr$^{3+}$ and Eu$^{3+}$ operating wavelengths, i.e 606 nm and 580 nm respectively in YSO. Because of the high frequency noise generated by the dye jet, it is a challenging task to achieve the sub-kHz linewidth sources that are needed to benefit from the long optical coherence lifetime. This limits the realization to laboratory proof-of-principle. Although some groups successfully built such laser sources [18,19], it is worth devising an alternative approach.

The thulium rare earth ion actually gathers together many advantages that make it specially attractive. First, its 793 nm wavelength falls within reach of easily stabilized semiconductor lasers, unlike Pr and Eu. Then, its $I = 1/2$ nuclear spin gives rise, under external magnetic field, to a simple straightforward 4-level optical system in which a Λ-system can be selected. Moreover, because of this simple structure, the sublevel splitting can be controlled easily by the external magnetic field. This crystal could therefore be used as an atomic quantum memory adapted to the bandwidth of existing quantum sources. For these reasons, thulium-doped materials are favorable alternatives to Pr- and Eu-doped compounds for quantum storage applications, with the triple advantage of a tractable wavelength, a simple level system and an adjustable ground state splitting.

Application of a magnetic field lifts the nuclear spin degeneracy, but this is not enough to obtain a Λ-system (cf Fig. 1). Indeed, optical excitation cannot flip a nuclear spin, as expressed by a selection rule on the nuclear spin projection $m_J$. However, coupling of electronic Zeeman effect and hyperfine interaction enhances the nuclear Zeeman effect and gives rise to nuclear gyromagnetic tensor anisotropy. If anisotropy is different in ground and upper electronic levels, the nuclear spin eigenvectors also differ in those two levels. As a con-
sequence the nuclear spin selection rule is relaxed. In YAG crystal, Tm ions are doped into low symmetry sites (D2) with a gyromagnetic tensor anisotropy that is much larger in the electronic ground state than in the excited state [17]. For an appropriate orientation of the applied magnetic field, the optical transition probability ratio along the two legs of the Λ can be optimized. In 2005, Guillot-Noël et al. theoretically showed that, for an adequate external field orientation, the branching ratio of the two transition probabilities reaches 0.24 in 2 crystalline sites out of the 3 that can be selected by laser beam polarization [20]. In 2006, we directly measured this optimal branching ratio to be 0.13 ± 0.02, proving that the nuclear spin selection rule is effectively relaxed [17, 21]. An adjustable Λ-system can now be built, involving both ground state sublevels coupled to one of the two excited state sublevels.

The magnetic field \( B \) being oriented to optimize the branching ratio, the nuclear Zeeman sensitivity can be measured with the help of hole burning spectroscopy. We find \( \Delta g/B = 36 \text{ MHz/T} \) and \( \Delta_e/B = 16 \text{ MHz/T} \), where \( \Delta_g \) and \( \Delta_e \) respectively stand for the ground and excited electronic state splittings. We also observe that, as depicted in Fig. 2, the side hole and antihole widths exceed the hole width at burning frequency, and vary linearly with the magnetic field amplitude with a slope of 0.99 MHz/T in the ground state, and 0.093 MHz/T in the excited state. We ascribe this broadening to spatial variations of the magnetic field and substitution site relative orientation. The external field orientation that optimizes the transition probability branching ratio is close to the direction of minimum splitting, and, accordingly, nearly orthogonal to the main component of gyromagnetic tensor. This is the reason why we measure a Zeeman sensitivity of 36 MHz/T, although the main gyromagnetic coefficient reaches 400 MHz/T in the ground state. As a consequence, a slight tilt of the field with respect to the site frame entails dramatic splitting variation. At 1 Tesla, a 1 mrad misorientation is enough to generate a 0.3 MHz splitting deviation in the ground state. At the moment it is not clear whether the observed inhomogeneous broadening is caused by site orientation disorder or by applied field orientational non-uniformity. The ground state width appears to be 10 times more sensitive to the magnetic field magnitude than the excited state width. Indeed the gyromagnetic tensor is much less anisotropic in the upper level than in the ground state. The ratio is consistent with an isotropic misorientation model.

We turn now to investigating the spin coherence with optical means. Initially the two ground state sublevels are equally populated and the optical transition frequency is distributed over a 25 GHz-wide inhomogeneously broadened absorption profile. Within this huge bandwidth, we select a narrow interval, smaller than \( \Delta_g - \Delta_e \), over which we prepare the ions for two-photon excitation by pumping them into a single sublevel. Optical pumping is accomplished by a sequence of 100 µs chirped pulses. Then we excite the ground state spin coherence with a 10 µs bichromatic laser pulse. The two frequency components \( \omega_1 \) and \( \omega_2 \), tuned to the Λ-system optical transitions, are separated by \( \Delta_g \) to achieve two-photon resonance. If all atoms remain phased together, one could monitor the spin coherence evolution by coherent forward Raman scattering: a long rectangular monochromatic pulse at frequency \( \omega_2 \) excites one transition of the Λ and converts part of the spin coherence into an optical coherence along the other transition. The resulting optical emission could be detected as a beatnote at frequency \( \Delta_g \) against the probe pulse. Unfortunately, the spin coherences evolve at different rates, according to the inhomogeneous broadening we observed in Fig. 2. This makes the beatnote vanish on the timescale of the excitation pulse duration. In order to recover the optical signature of the spin coherences, we resort to Raman echo procedure [22]. We apply an additional bichromatic pulse, resonant with the two-photon transition, at mid-time between initial excitation and final probing. This pulse reverses the spin coherence time evolution, so that, at the moment of probing, the spin coherences are phased together again and give rise to an optical emission. The Raman echo signal is observed by means of real-time Fast Fourier Transform.

The Raman echo is contaminated by two-pulse photon echo signals. The photon echo signal at frequency \( \omega_2 \) beats with the detection pulse, overlapping temporally and spectrally with...
the Raman echo. In previous Raman echo experiments \[^{[16]}\], photon echo was rejected by angular separation of the various signals. In the present work we have preferred a strictly collinear geometry that optimizes the spatial mode matching of the various beams. Then, to get rid of photon echo while maintaining an efficient driving of spin coherences, we detune the two frequencies of the rephasing pulse so that they are optically resonant with the other \(\Lambda\)-system of our 4-level system (cf Fig. 4).

The system is illuminated with an extended cavity diode laser (ECDL) operating at 793 nm, stabilized on a high-finesse Fabry-Perot cavity through a Pound-Drever-Hall servoloop down to 200 Hz over 10 ms \[^{[23]}\]. The laser is amplified with a semiconductor tapered amplifier (Toptica BoosTA). A polarizing cube split the beam. Each component is double-passed through an acousto-optic modulator (AOM) centered at 110 MHz (AA OptoElectronics). The two AOMs are driven by a dual-channel 1Gigasample/s waveform generator (Tektronix AWG520) that can provide arbitrary amplitude and phase shaping. In this experiment each channel feeds one frequency-shift at a time. After passing twice through the AOMs, the split beams come back to the cube where they merge into a fixed-direction single beam carrying the bichromatic excitation. A common polarization direction is given by a Glan prism. The recombined beam is finally coupled into a 2m-long single-mode fiber. The light polarization direction is adjusted with a half-wave plate to maximize the Rabi frequency. The laser frequency is then focused on the 5 mm-thick, 0.1 at.\% Tm\(^{3+}\):YAG sample cooled down to 1.7 K in an Oxford Spectromag cryostat. The magnetic field generated by superconducting coils is applied in the direction optimizing the branching ratio \[^{[21]}\]. The spot diameter on the crystal is 80 \(\mu\)m. The transmitted light is collected on an avalanche photodiode (Hamamatsu C5460 or C4777) protected from strong excitation damaging light pulses by a third acousto-optic modulator used as a shutter.

Raman echo experiments yield the spin coherence lifetime \(T_2\). The delay of the two bichromatic pulses is denoted \(T\). The Raman signal decays with \(T\) as \(e^{-2t/T_2}\). As an example, Fig. 4 shows the Raman signal decay for a \(\Delta_g = 41\) MHz ground state splitting. We measure the spin coherence lifetime for different ground state splittings in Tm\(^{3+}\):YAG. The low efficiency of the Raman echo process accounts for the low signal to noise ratio. Thanks to the double-pass setup, the two AOMs’ 50 MHz nominal bandwidth is extended to 100 MHz. However, 100 MHz is the upper boundary not for \(\Delta_g\) but for \(\Delta_g + \Delta_c \approx 1.4\Delta_g\), since the excitation pulses contain not only frequencies \(\omega_1 \) and \(\omega_2 = \omega_1 + \Delta_g\), but also \(\omega_1 + \Delta_c\) and \(\omega_1 - \Delta_g + \Delta_c\). At the limit of our equipment, we have yet been able to reach a ground state splitting \(\Delta_g\) up to 83 MHz. We show in Fig. 5 that the spin coherence lifetime does not significantly depend on the ground state splitting, remaining close to 300 \(\mu\)s over a 80 MHz range. The error bars correspond to the standard error deduced from a least-squares fit. We also measured the spin coherence lifetime in the excited state to be \(540 \pm 35\) \(\mu\)s for a 16.4 MHz excited state splitting with a similar Raman echo sequence. This lifetime is close to the population lifetime of the upper electronic state \[^{[24]}\]. With a population lifetime of 800 \(\mu\)s, the intrinsic spin coherence lifetime turns out to exceed 1 ms. The mechanisms responsible for spin decoherence in ground and excited states need to be unveiled by further studies.

Electromagnetically induced transparency (EIT) or controlled reversible inhomogeneous broadening (CRIB) \[^{[25]}\] experiments would be another step towards the demonstration of thulium-doped crystals as quantum memories. In materials such as ion-doped crystal, EIT might be regarded as a challenging operation. Indeed, overcoming the inhomogeneous linewidth by the coupling field Rabi frequency is known to be needed for efficient EIT \[^{[26]}\]. However, atoms far from optical resonance with the probe do not affect EIT, provided their ground state sublevels are equally populated. Therefore the effective inhomogeneous width to be considered essentially corresponds to the population-unbalanced atoms close to optical resonance. Those atoms cover a spectral interval much narrower than the inhomogeneously broadened absorp-
tion line. In either Pr$^{3+}$-, Eu$^{3+}$- or Tm$^{3+}$-doped crystals this feature is essential to discard non-resonant ions. In Eu$^{3+}$- and Pr$^{3+}$-doped crystals, resonant atoms tend to be pumped to the third sublevel by the driving fields and have to be repumped back to the selected $\Lambda$ system \cite{6,7} with the help of an auxiliary beam. This preparation procedure only affects atoms close to optical resonance. In Tm$^{3+}$-doped crystals this preparation step can probably be avoided since the ground state is split in two sublevels only.

Linear Stark shift should be very small in Tm$^{3+}$:YAG owing to the $D_2$ site symmetry. Therefore inhomogeneous broadening is difficult to reverse, which is not appropriate for CRIB. Alternatively, one can consider to dope Tm ions into host matrices exhibiting lower site symmetry such as Y$_2$O$_3$ or LiNbO$_3$. In these matrices, the Stark effect exists, allowing the building of an artificial inhomogeneous broadening by application of an electric field gradient. In LiNbO$_3$, the oscillator strength is found to be 50 times stronger than in YAG \cite{21}.

To conclude, for the first time we have optically excited, detected and characterized a nuclear spin coherence in a thulium-doped crystal. In addition to presenting a convenient absorption wavelength and a simple and adjustable $\Lambda$-type 3-level system, Tm$^{3+}$:YAG also offers long ground state spin coherence lifetimes, at splittings up to 80 MHz and probably much higher. This lifetime stability over a wide frequency range proves the crystal adequacy to existing quantum sources \cite{28,29}. The precise origin of the spin inhomogeneous broadening has still to be clarified. Given its specific properties, thulium can be considered as an excellent candidate for quantum storage in a solid, challenging the previously studied praseodymium or europium.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure5.png}
\caption{Measurement of the spin coherence lifetime for different ground state splittings, from 4 to 83 MHz. The error bars correspond to the standard error deduced from a least-squares fit.}
\end{figure}

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