Coherent Storage of Microwave Excitations in Rare Earth Nuclear Spins

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Interfaceing between various elements of a computer — from memory to processors to long range communication — will be as critical for quantum computers as it is for classical computers today. Paramagnetic rare earth doped crystals, such as Nd3+:Y2SiO5(YSO), are excellent candidates for such a quantum interface: they are known to exhibit long optical coherence lifetimes (for communication via optical photons), possess a nuclear spin (memory) and have in addition an electron spin that can offer hybrid coupling with superconducting qubits (processing). Here we study two of these three elements, demonstrating coherent storage and retrieval between electron and 145Nd nuclear spin states in Nd3+:YSO. We find nuclear spin coherence times can reach 9 ms at ~5 K, about two orders of magnitude longer than the electron spin coherence, while quantum state and process tomography of the storage/retrieval operation reveal an average state fidelity of 0.86. The times and fidelities are expected to further improve at lower temperatures and with more homogeneous radio-frequency excitation.

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Hybrid quantum systems composed of spin ensembles strongly coupled to superconducting resonators have recently emerged as a promising route for quantum memories operating in the microwave regime [1,2]. Such memories offer the possibility exploiting electron spin coherence times of up to seconds [3] as a resource for superconducting qubits, whose coherence times so far extend only to tens of microseconds [4]. Strong coupling has been observed between superconducting resonators and various paramagnetic impurities, including NV centres in diamond [5] and erbium ions in Y2SiO5 (YSO) and YAlO3 [6,7] leading to reversible coherent storage of (large numbers of) microwave photons within spin ensembles [8,9]. These paramagnetic impurities are often coupled to nuclear spins which can offer a further resource for storage — electronic spin coherences can be transferred to and from a nuclear spin [10–12], to access coherence times as long as hours [13].

The proposal to use solid-state spin ensembles as microwave quantum memories is in many ways inspired by results on using impurities in solid for optical quantum memories [14,15] where optical excitations are stored in rare earth (RE) nuclear spins [16–19]. Very long storage time can be expected, as nuclear spin coherence lifetimes in such materials extend up to 6 hours. Entanglement storage [20] and light-matter teleportation at telecom wavelength [21] have been demonstrated in Nd3+:Y2SiO5, for which optical coherence lifetimes of 90 µs have been measured [22,23].

Bringing together both optical and microwave strong coupling techniques on the same ensemble would enable a versatile quantum interface, connecting quantum memory, processing and communication and potentially allowing faithful conversion of microwave to optical photons [24,25]. However, hyperfine coherence lifetimes have so far only been studied for RE ions with an even number of f electrons and no electron spin [26,27]. It is therefore unknown whether nuclear spins could still provide a memory resource when the RE ions are paramagnetic, as required for coupling to microwave excitations. In this Letter we study a paramagnetic RE doped crystal Nd3+:Y2SiO5 and measure electron and nuclear spin coherence times of up to 100 µs and 9.2 ms, respectively. We further demonstrate coherence transfer between electron and nuclear spin degrees of freedom in the Nd3+:ion — quantum state and process tomography show transfer fidelities above the classical limit. These results suggest that quantum memories for microwave photons with access to long storage times are achievable in rare earth doped crystals.

Y2SiO5 (YSO) is a monoclinic crystal (C6h space group) with two crystallographic sites of C1 symmetry for Y3+ ions, which can be substituted by Nd3+ ions (Fig. 1(a)). Each site is divided in two classes related by a C2 symmetry along the crystal c axis. For magnetic fields parallel or perpendicular to the c axis, ions in the two classes are magnetically equivalent. Nd3+ has a [Xe]4f7 electronic configuration, with a 4H9/2 ground multiplet. In C1 symmetry, the crystal field (CF) splits the J multiplets into twofold degenerate levels. At low tem-
For some orientations of the magnetic field, weaker lines close to the $\mu$ about 3 pulses were 32 ns long and radio-frequency (rf) pulses were performed using an X-band (9.7 GHz) Bruker and D measurements were performed using an X-band (9.7 GHz) Bruker and D isotopically pure crystal boule of 0.001 at. % 145 Y, as well as 5 isotopes with zero nuclear spin. To reduce the concentration of ions not involved in the storage experiments and potentially causing dephasing, an isotopically pure crystal boule of 0.001 at. % 145 YSO was grown by the Czochralski method. Samples of about 1.5 mm$^3$ were cut with faces perpendicular to the b, D1 and D2 principal axes of the optical indicatrix. Experiments were performed using an X-band (9.7 GHz) Bruker electron spin resonance (ESR) spectrometer (Eleoxys 580) equipped with a helium cryostat. Microwave (mw) $\pi$ pulses were 32 ns long and radio-frequency (rf) pulses about 3 $\mu$s.

Figure 1(c) shows the field swept electron spin echo (ESE) spectrum obtained for a magnetic field oriented close to the D1 axis. The 16 intense lines correspond to the allowed ESR transitions ($\{\Delta m_I, \Delta m_S\} = \{0, \pm 1\}$) for the two magnetically inequivalent classes of one site. For some orientations of the magnetic field, weaker lines corresponding to $I = 0$ isotopes in the same site were observed. These results suggest that Nd$^{3+}$ ions preferentially occupy one of the Y$^{3+}$ crystallographic sites. The full linewidth at half maximum of the transition at 561.5 mT is 12 MHz, which is comparable to the narrowest linewidths measured in Er$^{3+}$:YSO, recently used to demonstrate strong coupling to a superconducting resonator [6]. All ENDOR and relaxation experiments below were performed at 561.5 mT. The Zeeman g and hyperfine A tensors were determined from CW spectra obtained by rotating the sample in planes containing the static magnetic field and perpendicular to the D1, D2 and b axes. A least squares fit to the ESR line positions gives the principal values of the g tensor: $g_x = 1.49$, $g_y = -0.98$, $g_z = -4.17$ with the Euler angles ($x \alpha \beta \gamma$ convention) relating the principal axes to the crystal axes D1, D2, b: $\alpha = 192^\circ$, $\beta = 39^\circ$ and $\gamma = 183^\circ$. In the same reference axes, the principal values of A and the corresponding Euler angles are: $A_x = 398$, $A_y = 0.1$, $A_z = 827$ MHz and $\alpha = 154^\circ$, $\beta = 34^\circ$ and $\gamma = 200^\circ$. As expected in low symmetry, the g and A tensors are highly anisotropic, but their principal axes are nearly parallel, as was observed for site 1 in Er$^{3+}$:YSO [56]

An electron-nuclear double resonance (ENDOR) spectrum was recorded (Fig. 1(d)), using a Davies ENDOR sequence with Tidy pulse [31][32]. The two ENDOR lines located at 201.7 and 202.8 MHz have Gaussian shapes with linewidths of 235 and 248 kHz respectively. Simulations confirm that these correspond to $+7/2: +7/2$ transitions in $m_I$, where the lower (higher) frequency line corresponds to the $m_S = +1/2$ ($m_S = -1/2$) transition. The coherence storage experiments described below involve the three transitions labeled $|1\rangle...|3\rangle$ as shown in Fig. 1(b).

The electron spin population relaxation time, $T_{1e}$, was measured by an inversion-recovery sequence as a function of temperature between 5 and 7 K. $T_{1e}$ increases with decreasing temperature from 0.1 to 30 ms (Fig. 2) and can be modeled above 5.5 K by an Orbach process with a CF level located 77 cm$^{-1}$ above the ground state, in reasonable agreement with the value of 88 cm$^{-1}$ deduced from optical measurements [53]. The electron spin coherence lifetime $T_{2e}$ was also studied in the same temperature range (Fig. 2), yielding stretched exponential decays with $T_{2e}$, in increasing from 28 to 106 $\mu$s with decreasing temperature. Stretched factors ranged between 1.2 and 1.5 below 6 K. We attribute the strong temperature dependence in $T_{2e}$ to the effect of spectral diffusion resulting from interactions with a bath of electrons spins undergoing spin-relaxation [54][55]. When the bath relaxation rate is much larger than the echo measurement scale, a stretch factor of $\approx 1.5$ indicates a Gaussian diffusion process [56]. Using this model and taking Nd$^{3+}$ ions themselves as the spin bath, $T_{2e}$ can be estimated from $T_{1e}$, the effective $g = 1.5$ and Nd$^{3+}$ concentration ($9.4 \times 10^{10}$ ions/cm$^3$), which gives $T_{2e} = 471$ $\mu$s. This is
Relaxation time (ms) | Temperature (K) | T2n
---|---|---
5.0 | 77 cm−1
6.2 | 5 K
6.6 | 7 K
5.4 | from 5 to 7 K.
5.8 | 2e component, an rf
7.0 | electron spin coherence to the

Before refocusing is complete, an rf π pulse to remove the effect of inhomogeneous broadening.

As this decay has the same origin as for the electron, we can simply relate $T_{2n}^\text{int} = \kappa T_{2e}$.

is about four times longer than the measured value and can be explained by the anisotropy of the g tensor, which increases the dipole-dipole interaction. Angular variation in the D1 − D2 plane showed that $T_{2e}$ is maximal in a region of about 5 degrees around D1 axis and decreases by a factor 2 at lower resonance fields.

Transfer between electron and nuclear spin coherences was performed using the sequence shown in Fig. 3(a) [10], which is fully compatible with the schemes designed for single photon operation [1, 2]. In our experiments, the memory input is a π/2 microwave pulse (consisting of $O(10^{17})$ photons). It creates an electron spin coherence on the $|1\rangle : |2\rangle$ transition, which is then refocused by a π pulse to remove the effect of inhomogeneous broadening. Before refocusing is complete, an rf π pulse on the $|2\rangle : |3\rangle$ transition transfers the coherence to $|1\rangle : |3\rangle$. At the time when this transition refocuses, a mw π pulse transfers the electron spin coherence to the $|2\rangle : |3\rangle$ NMR transition. To retrieve the coherent microwave signal from the nuclear spin ensemble, an rf π pulse refocuses the $|2\rangle : |3\rangle$ coherence, and then the sequence described above is applied in reverse order. A final mw π produces an electron spin echo, which is the output of the memory. This scheme allows extending storage times beyond $T_{2e}$, limited instead by the nuclear spin coherence time $T_{2n}$.

$T_{2n}$ was measured by monitoring the output echo amplitude as a function of $2\tau_n$ in the storage sequence (Fig. 3(a)). Echo decays were nearly exponential with maximal stretch factors of 1.25 and ranged from 184 μs at 7 K to 6 ms at 5 K (Fig. 2). $T_{2n}$ is bounded by $2T_{1e}$ when there is significant hyperfine coupling [10], and this limit is indeed observed for temperatures above 6 K. Below this temperature, some intrinsic nuclear spin decoherence mechanism is evident. We assume this intrinsic $T_{2n}$ follows the measured electron spin decoherence time $T_{2e}$, adjusted by some factor $\kappa$ to reflect the ratio of the effective g-factors for those ESR and NMR transitions: i.e. $1/T_{2n} = 1/(2T_{1e}) + 1/(\kappa T_{2e})$.

The fidelity of the coherent storage and retrieval into the $^{145}$Nd nuclear spin subspace was characterised using quantum state tomography and quantum process tomography at 6.5 K to avoid low repetition rates due to the long $T_{1e}$. Quantum state tomography is performed by measuring the qubit state in the Pauli basis ($\sigma_X, \sigma_Y, \sigma_Z$). Components $\sigma_X$ and $\sigma_Y$ are simply the real and imaginary part of the electron spin echo, while $\sigma_Z$ can be measured by an additional π/2 pulse immediately following the echo, to map $\sigma_Z$ onto $\sigma_X$ [10]. To obtain the overall process matrix of the electron-nuclear-electron spin transfer, density matrices are measured for the set of electron spin input states: $\pm |e\rangle X \pm |e\rangle Z$, and 1 (Fig. 3(b)). We are interested in obtaining the process matrix for the storage/retrieval operation itself, and so reference the output states against a simple two-pulse electron spin echo experiment with total duration equal to the time the coherent state resides in the electron spin degree of freedom, in the actual transfer sequence. In this way, losses related to electron spin relaxation, dephasing and state preparation are partly taken into account, but not errors related to the nuclear spin. The input and output states for the memory process are then linked by the relation, for a spin 1/2:

$$\epsilon(\rho_{\text{end}}) = \sum_{m,n=0}^3 \chi_{m,n} A_m \rho_{\text{start}} A_n^\dagger$$ (1)

where $\chi$ is the process matrix that is reconstructed, $A$ the operators from the Pauli basis ($I, \sigma_X, \sigma_Y, \sigma_Z$) and $\rho_{\text{start}}$ and $\rho_{\text{end}}$ the input and output density matrices (for a particular electron spin initial state) [37, 38].

We measure an average state fidelity (where $F_{\text{state}} = \text{Tr}(\sqrt{\rho_{\text{end}}} \sqrt{\rho_{\text{start}}})^2$) of $F_{\text{state}} = 0.86$, compared to what we would expect for an ideal memory, well above the classical limit of 2/3. The computed process matrix $\chi$ is shown in Fig. 3(c) and we find a process fidelity $F_p = \text{Tr}(\chi \chi_{\text{ideal}}) = 0.63$, where $\chi_{\text{ideal}}$ has just the identity component. Typically, average state and process fidelities are related by $F_{\text{state}} = (2F_p + 1)/3 = 0.75$ for a pure spin-1/2 [39]. However, preparation and measurements
are realized here on the electron spin, but conditional on a particular nuclear spin state, and so the reconstructed states do not span the full electron spin-1/2 state space. The reconstructed \( \chi \) process matrix was well simulated using a Linblad master equation and taking into account electron and nuclear spin relaxation rates \( (T_{1e}, T_{2e} \text{ and } T_{2n}) \), as well as pulse inhomogeneities (Fig. 3(c)). The latter were determined from fits to measured electron and nuclear spin Rabi oscillations (Fig. 4). The main process errors can be assigned to two particular contributions: first, the low fidelity of the rf pulses results in the large errors can be assigned to two particular contributions: first, the low fidelity of the rf pulses results in the large errors can be assigned to two particular contributions: first, the low fidelity of the rf pulses results in the large errors can be assigned to two particular contributions: first, the low fidelity of the rf pulses results in the large errors. The second contribution is pure dephasing, as evidenced by the \([\sigma_z, \sigma_z]\) component in \( \chi \), and is due to electron coherence decay during the application of the rf pulse. This could be significantly improved by lowering the temperature to increase \( T_{2e} \) (Fig. 2).

In conclusion, we have shown that microwave excitations can be stored into a nuclear spin coherence in a topically pure rare earth doped crystal, \(^{145}\text{Nd}^{3+} \cdot \text{Y}_2\text{SiO}_5\). Storage times, determined by the nuclear coherence lifetime, can reach 9.2 ms, about two orders of magnitude longer than the electron spin \( T_{2e} \) and the best superconducting qubit coherence times. Furthermore, these storage times could be significantly increased by dynamical decoupling techniques \(^\text{10}\). Given their long optical coherence lifetimes, our results show that paramagnetic rare earth doped crystals could be used as long lived quantum memories to interface superconducting qubits with both microwave and optical qubits.

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