Stopped light with storage times greater than one second using EIT in a solid

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Abstract

We report on the demonstration of light storage for times greater than a second in praseodymium doped Y$_2$SiO$_5$ using electromagnetically induced transparency. The long storage times were enabled by the long coherence times possible for the hyperfine transitions in this material. The use of a solid state system also enabled operation with the probe and coupling beam counter propagating, allowing easy separation of the two beams. The efficiency of the storage was low because of the low optical thickness of the sample, as is discussed this deficiency should be easy to rectify.

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Some of the most significant advances in quantum information processing have been made using quantum optics-based techniques. For example, working practical quantum cryptosystems already exist and there have been demonstrations of linear optics quantum computing [1], quantum teleportation, quantum non-demolition measurements [2], quantum feedback and control [3]. To proceed further it is necessary to have devices such as single photon sources, quantum memories and quantum repeaters, where quantum information is exchanged in a controlled fashion between light fields and material systems. It has been proposed that both the required control and strong coupling can be readily achieved using an ensemble approach, where the light field interacts with a large number of identical atoms. Such a ensemble based approaches now exist for single photon sources [4], ‘cat’ state sources [5], quantum memories [6, 7, 8] and quantum repeaters [4]. Experiments have demonstrated heralded single photon sources [9, 10] and the mapping of quantum information on a light field onto spin states of an atomic ensemble [11]. Experiments using electromagnetic induced transparency have demonstrated the storage and recall of optical pulses [12, 13].

The quantum systems used for these ensemble based demonstrations have almost exclusively been atomic vapors. An issue with these demonstrations is that even for a laser cooled ensembles, the atomic motion impacts on the devices’ performance. Ensembles of solid-state optical centers provide an alternative to atomic systems where the relative motion is zero. In this paper we investigate the use of solid-state system for ensemble based quantum optics and highlight its usefulness by stopping a light pulse using electromagnetically induced transparency (EIT). Unlike an earlier experiment [14], the current demonstration highlights for the first time two advantages of using optically active solid state centers: a one thousand fold increase in storage time and the ability to operate with a less restrictive beam geometry.

When storing light using EIT characteristics of the field are recorded as a spin wave in the ensemble. The storage time is determined by the coherence times of the hyperfine transitions. In principle coherence times for hyperfine transitions in atomic systems can be very long and many minutes have been measured in ion traps [15]. However, these long coherence times in large ensembles suitable for EIT have not been achieved. Transit time broadening in vapor cells and magnetic inhomogeneity in trapped systems mean that the longest that light has been stored atomic systems is a few milliseconds. In contrast, in earlier work we have demonstrated techniques to obtain hyperfine coherence times of tens of seconds in Pr:Y2SiO5 [16, 17]. Here we show it is possible to utilize these long coherence
times to stop light for similar lengths of time.

EIT is sensitive to atomic movement, with the spin wave being scrambled once the atoms have moved significantly compared to the wavevector mismatch between the probe and the coupling beams. To minimize this wavevector mismatch experiments in atomic systems typically operate with the beams co-propagating. Because the probe and coupling beams are close in frequency, in this configuration, the wavevector mismatch is typically less than 1 cm$^{-1}$. A consequence of this co-propagating operation is that it is difficult to separate the probe and the coupling beam. In a solid-state system, where the optical centers are locked in a crystal lattice, co-propagating operation is not required, in the present work the probe and the coupling beams are counter-propagating. With counter-propagating beams it is easier to separate the probe and coupling beam whilst maintaining optimum overlap.

The experimental setup is shown in FIG. 1. Because of the narrow 2500 Hz optical homogeneous linewidth of the $^3$H$_4\rightarrow^1$D$_2$ transition in Pr$^{3+}$:Y$_2$SiO$_5$ a highly frequency stabilized dye laser was required for the experiment not to be limited by laser jitter. The laser used was a modified Coherent 699 dye laser with a linewidth 200 Hz over 1 second time scales. The laser output was split into two beams, one of which was frequency shifted and gated by two AOMs and used as the probe beam. The other beam was frequency shifted and gated using a double pass AOM setup. This beam was used for the coupling and repumping fields. This coupling/repumping beams was aligned on a beam-splitter to go through the sample counter-propagating with the probe. The spare port of this right-most beam splitter was used to combine a local oscillator beam with the transmitted probe beam, enabling the heterodyne detection of the signal.

The sample used was the same as that used in reference [17] and consisted of 0.05% Praseodymium doped in Y$_2$SiO$_5$. It was 4 mm thick along the direction of light propagation. The sample was mounted in a bath liquid helium cryostat. Three orthogonal super-conducting magnets were used to apply a DC magnetic field to the sample and a six turn rf coil was used to apply a rf field.

The dominant dephasing mechanism for the hyperfine states of the Pr$^{3+}$ ions is random Zeeman shifting due to fluctuating magnetic fields from the yttrium nuclei. Dramatic increases in coherence times can be achieved by operating at a magnetic field where the transition frequency is insensitive to magnetic field changes to first order [16]. The magnetic field required is 78 mT in an orientation described in Ref. [16]. Once the magnetic field
FIG. 1: The experimental setup. What is not shown is a beam picked of the laser was put in the remaining port of the right most beam-splitter. This enabled heterodyne detection of the probe beam that was transmitted through the sample.

is obtained the remaining fluctuations have reasonably long correlation times. This situation enables the effective use of dynamic decoherence control (DDC) techniques and coherence times in excess of half a minute have been demonstrated.

An energy level diagram showing the transitions driven during the experiments is shown in Fig. 2. While the optical inhomogeneous line widths is a few GHz. The narrow homogeneous linewidth (of order 1 kHz) and long hyperfine population lifetimes (of order 1 minute) enabled the experiment to be carried out on an ensemble with a much smaller range of optical frequencies. At the beginning of each shot a sequence of the five optical frequencies (labelled “R” in Fig. 2) was applied repeatedly. The repump frequencies were applied sequentially rather that all at once to avoid the possibility of darks states and nonlinear mixing of the different frequencies in the AOM. The gap in time between the repumping and each experimental shot was long enough to ensure that ions had no remaining optical coherence. This repumping procedure prepared an ensemble of ions in the desired hyperfine state and gives a narrow adsorption with an inhomogeneous width of 100 kHz when measured by sweeping a week probe in frequency (line given by dots in Fig. 3). When the coupling beam was applied a narrow transparency was obtained in the absorption of the weak probe (solid trace in Fig. 3).

The repumping beams were applied after each shot and the 300 kHz span shown was swept in 4 ms. The transmitted probe beam was detected as a heterodyne beat signal and the bandwidth of the RF detector was comparable to 300 kHz, the extra noise at each end of the spectrum came from dividing out this frequency response. For coupling intensities above 1 mW the EIT was observed to depend linearly on the amplitude of the coupling.
FIG. 2: Energy level diagram showing transitions driven as part of the experiment. The experiment was carried out on the zero phonon line of the $^3\text{H}_4 \rightarrow ^1\text{D}_2$ optical transition. The hyperfine levels are shown and these are due to the $5/2$ spin of the praseodymium nuclei. In the presence of a magnetic field these are linear combinations of the zero field states. The probe, coupling and repump beams are labelled P, C and R respectively.

FIG. 3: Transmission of a weak probe, 10 $\mu$W, as its frequency is swept through resonance with the prepared ensemble. The solid line was taken with a 1 mW coupling beam on and the dotted line with the coupling off.
FIG. 4: On the left is the time sequence used in the stopped light experiments (a) with simple rephasing of the inhomogeneous broadening of the spin transitions and (b) with “bang-bang” dynamic decoherence control. In (a) two rephasing pulses are used placed 1/4 and 3/4 of the way through the storage time. In (b) N rephasing pulses were used (N even). The first rephasing pulse was applied 2 ms after the light was stored, the pulses were separated by 4 ms and the last pulse was applied 2 ms before the light was recalled. The rephasing pulses lasted 22 µs. On the right is the size of the recalled pulse as a function of time. The faster decay was acquired using simple rephasing of the ground state spin coherence (a). The slower decay was acquired using “bang-bang” decoherence control (b). The inset shows the energy of the recalled pulse as a function of the energy of the input pulse, the probe pulse length was 20 µs and the delay held constant at 100 ms.

beam. The limiting EIT width at low intensity was 10 kHz, corresponding to the hyperfine inhomogeneous linewidth. Below 1 mW the EIT transmission decreased with decreasing coupling intensity.

It can be seen from FIG. 3 that the peak absorption of our ensemble is only about 15% and, as is discussed below, this limits the efficiency of the storing process.

The time sequence for the light storage demonstration is shown on the left of FIG. 4. A 20 µs long probe pulse was applied and then the 10 mW coupling beam was turned off to transfer the optical coherence onto the spin transition. As in the previous solid state stopped light experiment RF rephasing pulses were used to rephase the inhomogeneous broadening in the spin transition. Although one RF rephasing pulse is enough to rephase the
spin-wave it also flips the spin-wave’s direction. Therefore when not using co-propagating beams, as is the case here, it is necessary to use an even number of rephasing pulses.

The size of the pulse of light recalled as a function of delay can be shown with and without dynamic decoherence control (DDC) and the results are shown in FIG. 4. The decay constants for the stored signal output were 0.35 seconds without DDC and 2.3 seconds with DDC. These decay rates were comparable to measurements of $T_2$ made using the same method as Fraval et al. The difference between the present measurements of $T_2$ and those obtained by Fraval et al. is attributed to not having tuned the magnetic field as carefully as was achieved by Fraval et al.

Shown in the inset of FIG. 4 is the intensity of the output pulse as the intensity of the input probe pulse is varied. From the graph it can be seen that the storage process is linear at low powers and starts to saturate at higher powers once the input pulse becomes a significant fraction of a $\pi/2$ pulse. This demonstration of linearity is important. Previous solid state experiments have been restricted by laser frequency jitter to using probe pulses with areas greater than $\pi$. At such high powers effects such as self induced transparency (SIT) cannot be ignored.

While the effect was linear and scaled to low powers, the efficiency was low, of the order of 1%. This in part can be improved with better timing and shaping of the probe and coupling waveforms. However the main reason for the low efficiency is the low optical absorption at the probe frequency and the accompanying modest group delay.

The sample used for this experiment was only 4 mm thick, longer samples as well as multi-pass cells and cavities are simple means to increase the optical absorption. Preliminary measurements on a samples with a range of praseodymium concentrations suggest that at least two or three fold increases in the optical thickness can be achieved by increasing the concentration without significantly increasing the inhomogeneous broadening of the hyperfine transition.

As it is a goal of this line of research to store and retrieve quantum mechanical states it worthwhile to consider the effect that rephasing pulses would have on few photon states stored in the hyperfine coherences. It has been asserted in a theoretical investigation of quantum information storage in the solid state that one would not be able to apply the RF $\pi$ pulses with sufficient accuracy. This is not the view of the authors of this paper. In Ref. it was assumed that the $\pi$ pulse would have to be applied with an accuracy close
to 1 part in $N$ (where $N$ is the number of atoms) in order that the few photon pulse not be swamped by light caused by inaccuracies of the $\pi$ pulse. However this light will be emitted randomly rather than in the very precise spatio-temporal mode of the output pulse. This should enable the output pulse to be easily separated from the background with very high efficiency.

In conclusion, we have demonstrate stopped light in Pr:Y$_2$SiO$_5$ for time scales of several seconds which is three orders of magnitude longer than any obtained previously. Based on previous measurements of $T_2$ it should be possible to extend this storage time by at least one more order of magnitude.

For the first time stopped light has been demonstrated in a solid with the coupling and probe beams counter propagating. This configuration is desirable as it allows easy separation of the two beams. However, it is only practical if the atoms movement during the storage time is small compared to the optical wavelength. Even for ultra-cold systems this places significant limits storage time. In a solid where the atoms are locked into position this isn’t a problem.

The efficiency of the storage process required for a quantum memory should be obtainable by increasing the density of the dopant ions and by increasing the interaction length.

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