Coherence rephasing combined with spin-wave storage using chirped control pulses

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Photon-echo based optical quantum memory schemes often employ intermediate steps to transform optical coherences to spin coherences for longer storage times. We analyze a scheme that uses three identical chirped control pulses for coherence rephasing in an inhomogeneously broadened ensemble of three-level Λ systems. The pulses induce a cyclic permutation of the atomic populations in the adiabatic regime. Optical coherences created by a signal pulse are stored as spin coherences at an intermediate time interval, and are rephased for echo emission when the ensemble is returned to the initial state. Echo emission during a possible partial rephasing when the medium is inverted can be suppressed with an appropriate choice of control pulse wave vectors. We demonstrate that the scheme works in an optically dense ensemble, despite control pulse distortions during propagation. It integrates conveniently the spin-wave storage step into memory schemes based on a second rephasing of the atomic coherences.

I. INTRODUCTION

Constructing optical quantum memories is of paramount importance for several applications in optical quantum information processing [1–3], e.g., for building quantum repeaters [4,5], or for linear optical quantum computing [6,7]. It is thus not surprising that devising and building such memories, devices with the capability to store and faithfully retrieve the quantum information contained in weak (few- or single-photon) light pulses, is currently a very lively field of research. A wide class of potential memory schemes use an inhomogeneously broadened atomic ensemble as a storage medium. The information carried by the amplitude and phase of the signal is mapped to atomic coherences as it is absorbed. These coherences promptly dephase, so a coherent optical response of the ensemble is prevented. However, if the dephased coherences can be rephased at a later time, an echo of the signal may be emitted—optical quantum information can be retrieved.

Numerous schemes of varying complexity have been proposed and demonstrated for rephasing the atomic coherences of the storage medium. For example, controlled reversible inhomogeneous broadening (CRIB) [2,8–10] and gradient echo memory (GEM) [11,12] schemes operate by broadening an initially narrow absorption line artificially with an inhomogeneous magnetic or electric field. Reversing the field also reverses the phase evolution of atomic coherences, so dephasing can be reversed. Atomic frequency combs (AFC) [13,14] involve preparing an absorption feature in the form of narrow, equidistant peaks. Large bandwidth signals are absorbed by atoms in multiple peaks, which first dephase, but later rephase spontaneously due to the discrete nature of the frequencies. To extend storage times and achieve on-demand retrieval, optical coherences in AFCs can be transferred to long-lived spin coherences between metastable ground states using strong control pulses [15,16]. These schemes involve a laborious preparation of the storage medium prior to signal absorption, but were demonstrated to function even at the single-photon level [17–20].

An earlier proposal [21] to store single-photon light pulses in inhomogeneously broadened ensembles was based on classical photon echos [22,23], which rely on strong control pulses for coherence rephasing. The simplest schemes of this type were shown to suffer from two major difficulties. The first one is noise from the inverted medium at the time of echo emission, which is incompatible with quantum information retrieval [24,25]. The second one is the distortion of short, intense π pulses—which are traditionally used as control pulses—while propagating in the optically dense medium [24,26]. The first one can be remedied by silencing the primary echo that would be emitted after the first control pulse and employing a second one to invert the medium again. The coherences can then be rephased a second time and a secondary echo will be emitted from an uninvited medium. One protocol, termed revival of silenced echo (ROSE) [27], can be realized by choosing the propagation direction of the control pulses such that the spatial modulation of the rephased coherences does not fulfill the phase-matching condition after the first pulse. The second flaw can be remedied by using frequency-chirped pulses that drive adiabatic passage (AP) between the atomic states as control pulses. With these improvements, traditional photon echos are compatible with few-photon signal storage [28] and are functional directly on telecom wavelengths [29].

Adiabatic passage driven by chirped pulses has been applied in a wide variety of fields for decades now [30,31]. Quite recently, AP was also employed successfully in various quantum memory applications, even though its use in such schemes is a somewhat subtle affair. Contrary to most applications of AP where only a robust population transfer is required, in quantum memory schemes the phase that the AP process imprints onto the atomic coherences is also very important. While manipulating the atomic populations, the overall phase associated with the process must also be essentially constant across the whole ensemble. One can show, however, that when two consecutive chirped pulses with identical amplitude and phase dependence are used to invert an ensemble of two-level atoms twice, the phases associated with each of the two AP processes cancel such that the overall phase will be the same for...
all atoms [16,27,32]. For this reason, AP by two chirped pulses can be used for implementing spin-wave storage in AFC memories [16], for spin-coherence rephasing in EIT based quantum memories [33,34], and for optical coherence rephasing to implement the ROSE scheme in two-level atoms [27–29].

The first advantage of AP in these schemes is, as in almost all other applications, that the precise parameters of the control pulses are not important; AP is robust with respect to parameter changes. The second one is that coherence rephasing can be realized with much smaller peak intensities than when short π pulses are used. This is especially important in solid-state media where the damage threshold of the crystal must not be exceeded. The third advantage of AP is its ability to function in optically dense ensembles. Population transfer and coherence rephasing induced by short π pulses are very fragile in an optically dense medium, because the control pulses are strongly distorted [24,26,35]. Population transfer and coherence rephasing induced by AP, on the other hand, is much more resistant to pulse distortion during propagation [35,36]. This latter is not at all trivial, because the two successive control pulses are distorted differently—one is absorbed by the medium, while the other one, propagating in the inverted medium, is amplified.

In this paper we consider the interaction between a series of chirped pulses and an inhomogeneously broadened, optically dense ensemble of three-level Λ systems. We show that a chirped pulse that interacts with both optical transitions of the system can realize an adiabatic rotation of the quantum states that results in a cyclic permutation of the atomic populations. Using three consecutive pulses, it is possible to regain the initial populations, and, at the same time, to rephase any optical coherences in the ensemble created by a signal prior to the control pulses. During one interval between the pulses, the information stored in optical coherences initially reside in spin coherences between the two lower levels. This sequence of control pulses thus integrates conveniently long-time spin-wave storage into the ROSE [27] protocol using control pulses from a single source. We investigate various aspects of the interaction relevant to coherence rephasing for photon-echo quantum memory applications. We consider the effect of the spectral width of the atomic ensemble relative to the full bandwidth of the control pulses. We discuss how echo emission during a possible partial rephasing while the ensemble is inverted can be suppressed by spatial phase mismatching as in the original ROSE scheme. Furthermore, we consider the question of control pulse propagation in the ensemble and identify the conditions under which the present scheme can rephase coherences in an optically dense sample. Finally, we discuss some constraints that the energy-level spacings of a material used for the realization of the scheme must fulfill and mention a specific example that does so.

II. PERMUTATION OF ATOMIC POPULATIONS WITH A CHIRPED PULSE

First, we study the effect of a frequency-chirped laser pulse on a single atom. It has three relevant energy eigenstates in a Λ configuration (Fig. 1); the frequency of the |1⟩ ↔ |2⟩ transition is \( \omega_{12} = \omega_0 + \Delta \), offset by \( \Delta \) from the line center \( \omega_0 \) of the inhomogeneously broadened ensemble. We assume that there is no broadening with respect to \( \omega_R \) and that decoherence effects can be neglected. The atomic Hamiltonian in a frame rotating with \( \omega_0 \) becomes \( H_a = \hbar \Delta \langle 2 | 2 \rangle + \hbar \omega_R | 3 \rangle \langle 3 | \). The pulse, polarized along \( \vec{e} \), interacts with both dipole allowed transitions, \( d_{12} = (1 | \vec{d} | 2 \rangle \langle 2 | \vec{d} | 2 \rangle ) \in R \)—the matrix elements are taken to be real, but not necessarily equal. We describe the atomic state with three probability amplitudes as \( | \psi \rangle = a | 1 \rangle + b | 2 \rangle + c | 3 \rangle \) and, using the usual dipole interaction Hamiltonian and the rotating-wave approximation, seek to derive the time evolution operator that propagates them from \( t = t_1 - T' \) just before the pulse to \( t = t_1 + T' \) just after.

Writing \( e(t, \vec{r}) = E(t, \vec{r}) e^{-i \omega_R t + i k \vec{r}} \), we can (locally) decompose the slowly varying complex field envelope of the pulse at the atom’s location \( E(t, \vec{r}) \) into a real amplitude and phase as \( A(t) e^{-i \Phi(t)} = d_{12} E(t, \vec{r}) \). Transforming to \( a(t) = a(t), b(t) = b(t) e^{i \Phi(t)}, \) and \( c(t) = c(t) \), and neglecting any decoherence, the relevant equation of motion will be

\[
\frac{\partial}{\partial t} \begin{pmatrix} a_r \\ b_r \\ c_r \end{pmatrix} = \frac{i}{2} \begin{pmatrix} A & 0 & 0 \\ 2 \delta & D \delta & -2 \omega_R \\ D \delta & -2 \omega_R & 0 \end{pmatrix} \begin{pmatrix} a_r \\ b_r \\ c_r \end{pmatrix},
\]

where we have introduced \( D = d_{12}/d_{31} \) and \( \delta(t) = \delta, \Phi(t) - \Delta \), the instantaneous detuning perceived by the atom. If \( A(t) \) and \( \delta(t) \) change sufficiently slowly, this equation can be solved by transforming to the reference frame of the adiabatic eigenstates at \( t_1 - T' \), neglecting any nonadiabatic transitions between these eigenstates and transforming back at \( t_1 + T' \).

The eigenvalue problem of Eq. (1) has been discussed at length in numerous papers, most notably in the context of simulated Raman adiabatic passage (STIRAP) [37–39]. Here we will simply summarize the relevant results; detailed formulas can be found, for example, in [40]. First, without losing generality we assume that \( \omega_R > 0 \). Then the matrix on the right-hand side has three distinct ordered eigenvalues for any \( \delta \), which we denote as \( \lambda^- < \lambda^0 < \lambda^+ \). In the limit when \( A \to 0 \), the eigenvalues and the corresponding eigenvectors \( \mu^{\pm,0} \) become

- For \( \delta < -\omega_R \), \( \lambda^+ \to 0 \), \( \mu^+ \to (1,0,0) \), \( \lambda^0 \to -\omega_R \), \( \mu^0 \to (0,1,0) \), \( \lambda^- \to \delta \), \( \mu^- \to (0,1,0) \).
- For \( \delta > 0 \), \( \lambda^+ \to \delta \), \( \mu^+ \to (1,0,0) \), \( \lambda^0 \to 0 \), \( \mu^0 \to (0,1,0) \), \( \lambda^- \to -\omega_R \), \( \mu^- \to (0,0,1) \).

If nonadiabatic transitions can be neglected, the time evolution operator from \( t_1 - T' \) to \( t_1 + T' \) in the adiabatic reference
frame is simply

$$
\hat{U}_{ad} = \begin{pmatrix}
\mathrm{e}^{i\lambda^+} & 0 & 0 \\
0 & \mathrm{e}^{i\lambda^0} & 0 \\
0 & 0 & \mathrm{e}^{i\lambda^-}
\end{pmatrix},
$$

where \( \lambda^{\pm,0} = \int_{t_1-T}^{t_1+T} \lambda^{\pm,0}(t) dt \)

is the integral of the adiabatic eigenvalues. For a pulse that is chirped from blue to red such that \( \delta(t_1 - T') > 0 \), \( \delta(t_1 + T') < -\omega_R \) (i.e., it becomes resonant with both atomic transitions), the time evolution operator in the original reference frame will be

$$
\hat{U} = \begin{pmatrix}
0 & \mathrm{e}^{i\lambda^+ + i\phi(t_1-T')} & 0 \\
0 & 0 & \mathrm{e}^{i\lambda^- - i\phi(t_1+T')} \\
0 & \mathrm{e}^{i\lambda^0} & 0
\end{pmatrix}. \quad (2)
$$

When the pulse is chirped from red to blue such that \( \delta(t_1 - T') < -\omega_R, \delta(t_1 + T') > 0 \), it becomes

$$
\hat{U} = \begin{pmatrix}
0 & 0 & \mathrm{e}^{i\lambda^0} \\
\mathrm{e}^{i\lambda^- - i\phi(t_1+T')} & 0 & 0 \\
0 & \mathrm{e}^{i\lambda^+ + i\phi(t_1-T')} & 0
\end{pmatrix}. \quad (3)
$$

The matrices in Eqs. (2) and (3) describe a cyclic permutation of the atomic populations with some additional phase factors. Pulses with opposite chirps permute the populations in an opposite sense. Figure 2 illustrates the process for a sech pulse with tanh chirp:

$$
\mathcal{A}(t) = A_0 \text{sech}(t/\tau_p), \\
\delta \phi(t) = \delta_0 + \mu \tau_p^{-1} \tanh(t/\tau_p).
$$

Figure 2(a) depicts the time evolution of the adiabatic eigenvalues for \( \mu < 0 \) (blue-to-red chirp), with the direction of population transfer between the three atomic states being illustrated in the inset with gray shading. The arrows that indicate the population transfer have been colored the same as the lines of the corresponding eigenvalues. Figure 2(b) is a similar plot for \( \mu > 0 \) (red-to-blue chirp). Because we have a single field and \( \omega_R \neq 0 \), we never have a two-photon resonance during the process—the pulse becomes resonant with the two single-photon transitions at different times. This means that we are in a regime distinctly different from that of STIRAP—we have no dark state. One can think of the process as two sequential adiabatic population transfers: first on the \([1] \leftrightarrow [2] \) transition, then on the \([2] \leftrightarrow [3] \) transition (for blue-to-red chirp). On the other hand, we do have an eigenstate that starts from \([1] \) and ends in \([3] \) (or the other way around). In certain parameter ranges, this contains only a small fraction of the excited state \([2] \), at any given time, so it may be quasidark [41].

The permutation of atomic populations is robust with respect to various parameter changes, as it is an adiabatic process. To show this, and in particular to quantify its efficiency, we have solved Eqs. (1) using a computer with various parameters, constructed the time evolution matrix, and calculated \( P_{\text{joint}} = |U_{12}U_{23}U_{31}|^2 \), the joint probability that there is a complete population transfer on all three transitions \([2] \leftrightarrow [1] \), \([1] \leftrightarrow [3] \), and \([3] \leftrightarrow [2] \) simultaneously. The pulse length \( \tau_p \) and the chirp parameter \( \mu \) were varied, while we had

**FIG. 2.** (Color online) Adiabatic permutation of atomic populations by a chirped pulse. (a) Solid lines: time evolution of the adiabatic eigenvalues for blue-to-red chirp; broken lines: eigenvalues for \( \mathcal{A} = 0 \) (diabatic lines). The direction of population transfer is indicated in the inset, where each arrow is the same color as the line of the eigenvalue belonging to the eigenstate that realizes the population transfer. (b) Eigenvalue evolution and direction of population transfer for red-to-blue chirp. The unit of frequency in the figure is \( \omega_R \); the pulse parameters are \( \tau_p = 10\omega_R^{-1}, A_0 = 2\omega_R, \delta_0 = -0.5\omega_R, \mu \tau_p^{-1} = -2/\mu \), and \( \Delta = 0, D = 1 \).

\( \Delta = 0, D = 1 \) and the amplitude of the sech pulse and the central detuning was always \( A_0 = 20/\tau_p \) and \( \delta_0 = -0.5\omega_R \) (i.e., the central frequency of the pulse was exactly halfway between \( \omega_{12} \) and \( \omega_{02} \)). The results are shown in Fig. 3, where (a) shows a contour plot of \( P_{\text{joint}} \) as a function of \( \tau_p \) and the chirp parameter \( \mu \tau_p^{-1} \), normalized by \( \omega_R^{-1} \) and \( \omega_R \), respectively.

**FIG. 3.** (Color online) (a) Contour plot of the joint probability that there is population transfer on all three atomic transitions \( P_{\text{joint}} = |U_{12}U_{23}U_{31}|^2 \) as a function of normalized pulse length \( \tau_p \omega_R \) and normalized chirp range \( \mu \tau_p^{-1}/\omega_R \). (b) Line plots of \( P_{\text{joint}} \) for three values of \( \mu \) as a function of \( \tau_p \).
The line plots in (b) show $P_{\text{gain}}$ for three specific values of $\mu_a$ as a function of $\tau_p$. It is clear from these figures that for the adiabatic permutation of populations to succeed, the transform limited bandwidth $\tau_p^{-1}$ must be much less than the spacing of the two lower levels, i.e., $\tau_p \omega_R \gg 1$. This condition ensures that the two optical transitions are traversed sequentially. Figure 3(b) shows that the joint probability is practically one for about $\tau_p \omega_R = 5$.

### III. COHERENCE REPHASING WITH A SERIES OF CHIRPED PULSES

#### A. Control pulses with negative chirp

We now consider a sequence of three consecutive chirped control pulses for coherence rephasing in an inhomogeneously broadened atomic ensemble. We have photon-echo based quantum memories in mind; the time line of the envisioned process is sketched in Fig. 4(a), while the transformation of the atomic states by the various pulses is sketched in Fig. 4(b). All atoms are in $|1\rangle$ initially; when a weak signal pulse, resonant with the $|1\rangle \leftrightarrow |2\rangle$ optical transition, is absorbed at $t_0$. The $a_j^\dagger b_j$ atomic coherences created by the signal dephase, so the overall ensemble polarization disappears. Three strong control pulses with negative (blue-to-red) chirp follow at $t_1$, $t_2$, and $t_3$, respectively. The first one at $t_1$ transforms the $a_j^\dagger b_j$ coherences into $a_j^\dagger c_j$ spin coherences between $|1\rangle$ and $|3\rangle$, where they can remain intact for a duration close to the spin-coherence time $T_{\text{storage}} \approx T_{\text{spin}}$. The second control pulse at $t_2$ transforms them into $c_j^\dagger b_j$ optical coherences on the $|3\rangle \leftrightarrow |2\rangle$ transition. There is population inversion in the ensemble at this point, so if rephasing should occur, echo emission must be suppressed as it will be too noisy for quantum memory applications [24].

Finally, the third control pulse at $t_3$ transforms coherences back to the $|1\rangle \leftrightarrow |2\rangle$ transition. If coherence rephasing succeeds, there will be a revival of the ensemble polarization and echo emission becomes possible at $t_4$.

To investigate whether coherence rephasing by the control pulses is indeed possible, we construct the time evolution operator that evolves the atomic states from $t = t_0 + T$, just after the signal pulse has been absorbed, to $t = t_1 - T$ just before the echo is emitted at $t_4$:

$$
\hat{U}(\Delta, \tilde{r}) = \hat{U}^F(t_4 - T, t_3 + T) \hat{U}_3(\Delta, \tilde{r}) \hat{U}^F(t_3 - T', t_2 + T') \times \hat{U}_2(\Delta, \tilde{r}) \hat{U}^F(t_2 - T', t_1 + T) \hat{U}_1(\Delta, \tilde{r}) \times \hat{U}^F(t_1 - T', t_0 + T).
$$

(5)

Here $\hat{U}_i(\Delta, \tilde{r})$ is the time evolution operator of the $i$th control pulse from $t = t_i - T'$ to $t = t_i + T'$, while $\hat{U}^F(t', t)$ is that for free evolution between the control pulses:

$$
\hat{U}^F(t', t) = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & e^{-i\Delta(t'-t)} & 0 & 0 \\
0 & 0 & e^{-i\omega_R(t'-t)} & 0 \\
0 & 0 & 0 & e^{-i\omega_R(t'-t)}
\end{pmatrix}.
$$

(6)

The $\hat{U}_i(\Delta, \tilde{r})$ depend on $\Delta$ and $\tilde{r}$, because the adiabatic eigenvalues $\lambda_{\pm,0}$ contain an explicit dependence on $\Delta$, and an implicit dependence on $\tilde{r}$ through $\mathcal{A}$ and $\Phi$ which vary in space as well.

For atoms with $\Delta$ such that all three control pulses permute the populations as described by Eq. (2), $\hat{U}(\Delta, \tilde{r})$ will be diagonal with

$$
[\hat{U}(\Delta, \tilde{r})]_{11} = e^{i\Lambda_2^0 + i\Lambda_1^0 + i\Lambda_3^1} 
\times e^{-i\Delta(t_1-t_2-2T')-i\omega_R(t_1-t_2-2T')},
$$

$$
[\hat{U}(\Delta, \tilde{r})]_{22} = e^{i\Lambda_1^0 + i\Lambda_2^0 + i\Lambda_1^0} 
\times e^{-i\Delta(t_2-t_3-t_4-2T')-i\omega_R(t_2-t_3-2T')},
$$

$$
[\hat{U}(\Delta, \tilde{r})]_{33} = e^{i\Lambda_1^0 + i\Lambda_2^0 + i\Lambda_1^0} 
\times e^{-i\Delta(t_3-t_4-2T')-i\omega_R(t_3-t_4-2T')}. 
$$

(7)

For brevity, we have not indicated the $\Delta$ and $\tilde{r}$ dependence of the $\Lambda_{\pm,0}$ and have dropped the unimportant $\Phi(t, \pm T)$ constant phase terms. The $a^*b$ coherence at $t = t_4 - T$ is then given in terms of its value at $t = t_0 + T$ as

$$(a^*b)' = a^*b \exp \left( i \left[ \Lambda_2^0 - \Lambda_1^0 + \Lambda_1^1 - \Lambda_2^1 + \Lambda_3^1 - \Lambda_2^0 \right] \right)
\times \exp(-i\Delta(t_1 - t_0 + t_4 - 2t_3 + t_2 - 2T))
\times \exp(-i\omega_R(t_1 - 2t_2 + t_3)).
$$

(8)

If the three control pulses are identical, the corresponding $\Lambda_{\pm,0}$'s are equal, so the argument of the first exponential vanishes for any $\Delta$. The third exponential describes a uniform phase shift for the whole ensemble, while the second one is $\exp(i2\Delta T)$ provided that the time intervals between the pulses fulfill

$$
t_1 - t_0 + t_4 - t_3 = t_3 - t_2. 
$$

(9)

Therefore, with three identical control pulses, the atomic coherences will be rephased at $t_4$—an echo of the signal can be emitted. Equation (9) also shows that rephasing does not depend on the duration $t_3 - t_1$ when the coherences generated by the signal reside in the $|1\rangle \leftrightarrow |3\rangle$ spin coherence.
The properties of coherence rephasing with three positively chirped pulses can be derived in an analogous manner from the time evolution matrices 3 and 6. The order in which the quantum states are traversed is now reversed compared to that shown on Fig. 4(b). The medium will be inverted after the first control pulse between $t_1$ and $t_2$, while spin-wave storage will take place between $t_2$ and $t_3$. The relevant formula for the evolution of the atomic coherences from $t = t_0 + T$ to $t = t_4 - T$ is

$$
(a^* b)' = a^* b \exp(i\lambda_0^0 - \lambda_0^1 + \lambda_3^1 - \lambda_3^0 - \lambda_2^- - \Lambda_1^-) \exp(-i\Delta [2t_1 - t_0 + t_2 - t_3 - t_2 - 2T]) \times 
\exp(i\omega_R(t_1 - 2t_2 + t_3 - t + T')).
$$

Again, for three identical control pulses the argument of the first exponential vanishes, while the second exponential gives $t_1 - t_0 + t_2 - t_3 = t_2 - t_1$ for the condition of rephasing. As before, we must also consider a possible rephasing in the inverted medium, this time after the first control pulse. The $|3\rangle \leftrightarrow |2\rangle$ coherence at $t_3 - T$ will be

$$
(c^* b) = ab^* \exp(i\lambda_0^0 - \lambda_0^1) \exp(i\Delta(t_1 - t_0 + t_1 - t'_2)) \times 
\exp(-i\omega_R(t_1 - 2t_2 + t_3 - T + T')).
$$

The second exponential will certainly disappear at some $t'_2 \in [t_2, t_3]$ because we need $t_1 - t_0 < t_2 - t_1$ to obtain the secondary rephasing. While the first exponential will not be zero, it may again be too weakly dependent on $\Delta$ to extinguish the primary echo fully.

### IV. Towards a Quantum Memory

#### A. Silencing the primary echo

As discussed for ROSE in two-level atoms [27,35], the primary echo in the inverted ensemble can be silenced with an appropriate choice of the control pulse wave vectors. If the spatial modulation of the revived ensemble polarization does not fulfill the phase-matching condition, a collective emission is not possible—coherence rephasing does not lead to echo emission. Assuming plane-wave control pulses and taking into account the $\sim \exp(i\vec{k}_1 \cdot \vec{r})$ fast spatial modulation of the $l$th amplitude, we can derive the spatial modulation of the coherences after each pulse. We can then obtain the following phase-matching conditions for the echo wave vectors:

- for negative chirp: $\vec{k}_e = 2\vec{k}_3 - \vec{k}_2 - \vec{k}_1 + \vec{k}_s$,
- for positive chirp: $\vec{k}_e = 3\vec{k}_3 - 2\vec{k}_2 + \vec{k}_1 + \vec{k}_s$. (10)

Here $\vec{k}_e$ is the wave vector of the signal field and $\vec{k}_e^{(i)}$ denotes the wave vector of the spatial modulation of the coherences after the $i$th control pulse. We assume that the wave vectors of the control pulses and the signal field are all approximately equal in magnitude $k_e = k_1 = k_3 = k_s$. (More precisely we assume that for any difference in the wave vectors $L\Delta k \ll \pi$, where $L$ is the spatial extent of the storage medium.) Echo emission is possible if there is a revival of the ensemble polarization due to (partial) rephasing and $\vec{k}_e^{(i)} = k_e$. The wave vector of the secondary echo we want emitted in both cases is $\vec{k}_e^{(3)}$, whereas $\vec{k}_e^{(2)}$ and $\vec{k}_e^{(1)}$ are the wave vectors of the primary echoes we want suppressed. From Eqs. (10) we can deduce the following:

- (i) For control pulses collinear with the signal, all propagating in the same direction $\vec{k}_1 = \vec{k}_2 = \vec{k}_3 = \pm \vec{k}_s$, we have a forward secondary echo $\vec{k}_e^{(3)} = \vec{k}_s$.

- (ii) If $\vec{k}_1 = \vec{k}_2 = \vec{k}_3 = -\vec{k}_s$, i.e., we have backward propagating control pulses, $\vec{k}_e^{(2)} = -3\vec{k}_s$ and $\vec{k}_e^{(1)} = -3\vec{k}_s$, so the primary echo is silenced.

- (iii) If we are not restricted to signal and control pulse propagation along a single direction, we can obtain a backward propagating secondary echo $\vec{k}_e^{(3)} = -\vec{k}_s$ with the setups sketched in Fig. 5.

Note that if $\omega_R < 0$ the formulas and figures for positive and negative chirps must be interchanged.

#### B. Ensemble spectral width and chirp range

In Secs. II and III we have seen that coherence rephasing in an ensemble requires the frequency sweep of the control pulses to encompass both optical transitions ($\omega_{22}$ and $\omega_{32}$) of the atoms to be rephased. At the same time, the distribution of $\Delta$’s must be wide enough to absorb all frequency components of the
signal—the ensemble spectral width \( \sigma_A \) must be greater than the signal bandwidth. Let us assume that the signal spectrum is bounded by \( \omega_0 \pm \sigma_s \), i.e., this is the spectral range that the control pulses must rephase. Assuming pulses as in Eqs. (4), the range of \( \Delta' \)'s where both optical resonances are crossed is \( \Delta' \in [\Delta_0 - \mu \tau_p^{-1} + \omega_R, \Delta_0 + \mu \tau_p^{-1}] \), while the spectral region properly rephased is expected to be slightly narrower due to a transition region of width \( \sim \tau_p^{-1} \) at the edges of this interval where nonadiabatic transitions are possible. Centering the interval on \( \Delta = 0 \) with \( \delta_0 = -\omega_R/2 \), the condition for \( \tau_p \) and \( \mu \) of the control pulses becomes

\[
\Delta_{\text{max}} = |\mu \tau_p^{-1} - \omega_R/2| > \sigma_s.
\]

Note that because we assume that \( |3 \rangle \) is empty initially and that the signal field is very weak, we do not need \( \omega_R > \sigma_s \)—the signal bandwidth can be larger than the separation of the two lower states.

After determining the minimal chirp range, we must also compare \( \Delta_{\text{max}} \) to the ensemble spectral width \( \sigma_A \). This is an important question, because the unmanipulated absorption lines of atomic systems used in optical quantum memory experiments are often much wider than the spectrum of the control pulses. For example, in rare-earth ion doped crystals that are popular candidates for such devices, very typical orders of magnitude are \( \sim 1 \) GHz for the inhomogeneous broadening, \( \sim 10 \) MHz for hyperfine splittings of the lower states, and \( \sim 1 \) MHz for signal bandwidths [18,20,28]. In these cases, quantum memory schemes usually work only if the absorption line is first tailored to a sufficiently narrow range (for example, using optical pumping). In the original ROSE scheme, a considerable number of atoms may be left excited after the second control pulse if \( \pi \) pulses are employed for control and an unmanipulated ensemble is used as a storage medium. These are then a source of spontaneous noise during signal retrieval. However, if chirped pulses are used for coherence rephasing, the number of atoms left excited in the spectrally wide ensemble can be orders of magnitude smaller [27,35].

Figure 6 depicts the effect of the control pulses on a wide ensemble. It shows the final populations of the atomic states after the three control pulses with parameters \( \tau_p = 1 \mu s, \omega_R = 10 \text{ MHz}, \mu \tau_p^{-1} = -30 \text{ MHz}, \) and \( A_0 = 16 \text{ MHz}. \) With these parameters, atoms are rephased roughly for \( \Delta \in [-23 \text{ MHz}, 23 \text{ MHz}] \), the spectral range where they are returned to \( |1 \rangle \) finally (the central part of the broken blue line on Fig. 6). Apart from the transition regions where some atoms are left partially in \( |2 \rangle \) due to nonadiabatic transitions, there is also a wide plateau where there is full excitation after the control pulses. This happens because for atoms with \( \Delta \in [-35 \text{ MHz}, -25 \text{ MHz}] \) the control pulses never become resonant with \( \omega_{32} \), so these atoms behave like two-level atoms. They are inverted three times by the control pulses and end up in \( |2 \rangle \) after the third one (solid red line on Fig. 6). So, in a spectrally wide ensemble, there is an \( \sim \omega_R \) wide region where atoms are left excited after the control pulses. This region will be a source of spontaneous noise at the time of signal retrieval. Because \( \omega_R \gg \tau_p^{-1} \), the situation is much less favorable than for two-level atoms, where the regions at the edge of the control pulse bandwidth with remanent excitation were \( \sim \tau_p^{-1} \) in width [35]. Thus, to use the current rephasing scheme for quantum memory applications, it is necessary to tailor the absorption line to a width such that both atomic resonances are encompassed by the control pulses. However, this does not mean that the optical depth of the storage medium will be reduced at the signal frequency. All atoms with resonance frequencies within the signal spectrum are used and rephased by the control pulses. By contrast, schemes like CRIB and GEM must first tailor an absorption line which is narrow compared to the signal bandwidth, leading to a loss of optical depth.

C. Rephasing coherences in an optically dense ensemble

The next question to be considered is the ability of the chirped control pulses to rephase an optically dense ensemble. In photon-echo based memories we need optically dense samples to absorb the signal—but these also distort the control pulses propagating within them considerably. Moreover, consecutive control pulses will be distorted in different ways. A pulse that excites the ensemble is absorbed, one that moves atoms back to one of the ground states is amplified, while a pulse that moves atomic populations between the two stable states can propagate without considerable distortion to great optical depths [42]. Therefore, even if the control pulses can rephase the ensemble at the entry, their ability to do so further into the medium is a question. In previous investigations, it was
found that short $\pi$ pulses are very fragile in optically dense ensembles of two-level atoms [24,26], but two consecutive chirped pulses can be used to rephase coherences to sufficient optical depths [35].

To answer this question, we have solved the relevant Maxwell-Bloch equation for the propagation of the control fields using a computer. To obtain a tractable problem, we have assumed that the control pulses are plane waves propagating in the same direction along the $z$ axis, i.e., we considered a one-dimensional problem only. In the slowly varying envelope approximation, the wave equation for the complex pulse amplitudes $\Omega_j(z,t) = E_j(z,t)d_{12}/\hbar$ will be

$$\left(\partial_z + \frac{1}{c} \partial_t\right) \Omega_j(z,t) = i \frac{\alpha_d}{\pi g(0)} \int [a^* (z,t,\Delta)b(z,t,\Delta) + Dc^* (z,t,\Delta)b(z,t,\Delta)] g(\Delta)d\Delta.$$  

(11)

Here $\alpha_d = \pi g(0)kN d_{12}^2/\epsilon_0 \hbar$ is the absorption constant of the $|1\rangle \leftrightarrow |2\rangle$ transition, $g(\Delta)$ is the spectral distribution function of the ensemble, and $N$ is the density. The integral on the right is the overall polarization of the ensemble at time $t$ and point $z$—we now have an ensemble extended in both $\Delta$ and $z$, so the probability amplitudes $a, b,$ and $c$ depend on these variables too. Equation (11) must be solved together with the optical Bloch equations

$$\partial_t \begin{pmatrix} a \\ b \\ c \end{pmatrix} = \frac{i}{\Delta_1} \begin{pmatrix} 0 & \Omega_j^* & 0 \\ \Omega_j & -2\Delta & D\Omega_j \\ 0 & D\Omega_j^* & -2\rho_0 \end{pmatrix} \begin{pmatrix} a \\ b \\ c \end{pmatrix}$$

(12)

for the atoms of the ensemble. [This is the same as Eq. (1) rewritten in terms of a complex field amplitude $\Omega_j.$] The equations were solved for three consecutive, copropagating chirped control pulses with blue-to-red chirp. Because we have optical quantum memories in mind, the excitation that signal absorption generates in the medium is completely negligible when considering control pulse propagation; we can assume all atoms to be in $|1\rangle$ initially.

Having obtained $\Omega_j(t,z)$ for all three pulses, we can readily calculate the time evolution matrices $U_j(\Delta,z)$ by solving Eqs. (12) again for a set of atoms with various $\Delta$’s and initial conditions. To investigate the extent of the domain where coherences are rephased, we define the rephasing factor $R(\Delta,z)$ using the auxiliary quantities $R_1(\Delta,z)$ and $R_2(\Delta,z)$:

$$R_1(\Delta,z) = [\hat{U}_1(\Delta,z)]_{21}[\hat{U}_2(\Delta,z)]_{12}[\hat{U}_3(\Delta,z)]_{13},$$

$$R_2(\Delta,z) = [\hat{U}_1(\Delta,z)]_{13}[\hat{U}_2(\Delta,z)]_{12}[\hat{U}_3(\Delta,z)]_{13},$$

as

$$R(\Delta,z) = R_1^*(\Delta,z)R_2(\Delta,z).$$

(13)

Comparing Eqs. (2), (7), and (8), one can see that $|R_1(\Delta,z)|^2$ is the probability that an atom at $z$ with frequency offset $\Delta$ in state $|1\rangle$ initially is transferred to $|2\rangle$, then to $|3\rangle$, and finally back to $|1\rangle$ by the control pulses, while $\arg[R_1(\Delta,z)]$ is the overall phase associated with this process. $|R_2(\Delta,z)|^2$ is the probability that an atom in $|2\rangle$ initially is moved to $|1\rangle$, then to $|3\rangle$, and finally back to $|2\rangle$, while $\arg[R_2(\Delta,z)]$ is the associated phase. For $\Delta$ and $z$ such that all three control pulses drive AP between the atomic states, $R(\Delta,z)$ is precisely the first exponential factor in Eq. (8) that contains the eigenvalue integrals $\Lambda_j^{+1/0}$. In general, $R(\Delta,z)$ is an overall factor whose magnitude gives the probability that all populations have been permuted three times as required, and also the associated phase factor, whether AP has taken place or not. Clearly, to be able to rephase the coherences in some domain of the atomic ensemble, we must have $|R(\Delta,z)|^2 = 1$ and $\arg[R(\Delta,z)] = \text{const}$ in an interval of $\Delta$ and $z$.

Note that we cannot simply construct the overall transfer matrix $\hat{U}(\Delta,z)$ and use its diagonal matrix elements to investigate coherence rephasing. $\hat{U}_{11} = R_1$ and $\hat{U}_{22} = R_2$ is true only if all three pulses drive AP, i.e., if all elements of the $\hat{U}_j$ associated with nonadiabatic transitions are zero.

The control fields $\Omega_j$ and the rephasing factor $R$ were first calculated for an atomic ensemble whose spectral distribution was taken to be a constant in the range $\Delta \in [-20 \text{ MHz}, 20 \text{ MHz}]$ and zero elsewhere. Control pulse parameters were the same as those used for the calculation depicted in Fig. 6, so this case amounts to tailoring the absorption line such that the pulses can perform the permutation of the atomic populations for the whole ensemble. Figure 7(a) shows the magnitude of the three amplitudes at an optical depth of $\alpha_dz = 5$. Clearly, the $\Omega_j$ are different here, even though they were identical at $\alpha_dz = 0$. $\Omega_1$, which transfers atoms from $|1\rangle$ to $|3\rangle$, is not really changed, $\Omega_2$, which excites atoms, is attenuated, while $\Omega_3$, which returns them to $|1\rangle$, is amplified. Figures 7(b) and 7(c) depict $|R(\Delta,z)|^2$ and $\arg[R(\Delta,z)]$ (in radians) as a function of the optical depth $\alpha_dz$ and the frequency offset $\Delta$. The black line in (b) marks $|R(\Delta,z)|^2 = 0.99$. The plots demonstrate that the control pulses can in fact rephase a considerable part of the ensemble.
in the previous case [see Fig. 7(a)]. Increasing the pulse to a width that allows permutation of the atomic populations
applications, the ensemble spectral width has to be tailored
rephasing scheme to be useful for optical quantum memory
amplitudes does not help; using more intensive pulses yields

![FIG. 8. (Color online) Control pulse propagation in an optically dense ensemble. (a) Magnitude of the control pulses at an optical depth of $\alpha_d z = 5$. (b) Contour plot of $|R(\Delta, z)|^2$ and (c) arg($R(\Delta, z)$) measured in radians. $g(\Delta) = g(0)$ for $\Delta \in [-50\text{ MHz}, 50\text{ MHz}]$; $g(\Delta) = 0$ elsewhere. Pulse parameters are the same as for Fig. 6.](image)

The magnitude of $R$ is very nearly 1 for the whole spectral width until about $\alpha_d z = 5$, and the phase difference is also small ($\lesssim \pm 10^{-1}$ rad) at $\alpha_d z = 5$ for a spectral width of about 14 MHz. This seems sufficient, as it would allow a signal of $\sim 10$ MHz bandwidth to be stored and emitted by the ensemble—an optical depth of $\alpha_d z = 5$ allows the absorption of 99.3% of the signal pulse energy. Note that for forward echos, which is the only possibility for control pulses propagating along a single direction, $\alpha_d z = 2$ is the ideal choice [10].

The $\Omega_j$’s and $R$ were also calculated for an atomic ensemble whose spectral width was wide with respect to the bandwidth of the control pulses [$g(\Delta) = \text{const}$ was assumed in the range $\Delta \in [-50\text{ MHz}, 50\text{ MHz}]$]. Figure 8(a) depicts the $|\Omega_j|’s$ at $\alpha_d z = 5$, while 8(b) and 8(c) depict $|R(\Delta, z)|^2$ and arg($R(\Delta, z)$) (in radians). Clearly, this time the ability of the same control pulses to rephase the coherences of the ensemble deteriorates much more quickly as they propagate. $|R(\Delta, z)|^2$ drops below 0.9 by about $\alpha_d z = 3$ everywhere, and the phase differences are also much greater than they were in the previous case. We attribute this loss of rephasing to pulse distortions that arise from the interaction of the control pulses with atoms for which $\omega_{32}$ is outside the control pulse spectrum, i.e., which behave as two-level atoms. The pulse amplitudes at $\alpha_d z = 5$ shown in Fig. 8(a) are much more distorted than in the previous case [see Fig. 7(a)]. Increasing the pulse amplitudes does not help; using more intensive pulses yields very similar results. Therefore, we conclude that for the current rephasing scheme to be useful for optical quantum memory applications, the ensemble spectral width has to be tailored to a width that allows permutation of the atomic populations for the entire ensemble. However, we stress again that this does not mean that we lose any optical depth at the signal frequency.

D. Material considerations

Finally, we consider some material properties that are required for the realization of our scheme. A number of quantum memory schemes have been demonstrated in rare-earth (RE) doped optical materials, where the hyperfine levels of the dopant ions offer the possibility of selecting an atomic system with a $\Lambda$ configuration for spin-wave storage (see, e.g., [15,20,43]). Apart from the various homogeneous linewidths, there is a special requirement in our case. Because the same control pulses are to become resonant with two optical transitions one after another, it is important that there be no other atomic transitions between the two used for realizing the $\Lambda$ linkage. As hyperfine transitions in RE doped crystals cannot usually be polarization selected, this effectively means that if the excited state has multiple hyperfine sublevels, their separation must be greater than that of the two ground-state hyperfine levels used.

One possible example of such a material is $^{153}\text{Eu}^{3+}:\text{Y}_2\text{SiO}_5$, whose hyperfine levels have been sketched in Fig. 9 on the left, taken from [44]. Making the correspondence $|1\rangle \leftrightarrow \pm (5/2g)$, $|3\rangle \leftrightarrow \pm (1/2g)$, and $|2\rangle \leftrightarrow \pm (5/2e)$, to obtain a $\Lambda$ system and denoting the frequency of the $\pm (3/2g) \leftrightarrow \pm (5/2e)$ transition by $\omega_{32}$, we have $\omega_{32} = \omega_{32} - 2\pi \times 90\text{ MHz}$, i.e., $\omega_{51} = 2\pi \times 90\text{ MHz}$. There are no optical transitions between $\omega_{32}$ and $\omega_{51}$; the two closest to them will be the $\pm (5/2g) \leftrightarrow \pm (5/2e)$ transition at $\omega_{32} + 2\pi \times 119\text{ MHz}$ and the $\pm (5/2g) \leftrightarrow \pm (3/2e)$ transition at $\omega_{32} - 2\pi \times 51\text{ MHz}$. This means that using control pulses of length $\sim 0.1\text{\mu s}$ and a chirp ranging from $\omega_{32} - 2\pi \times 130\text{ MHz}$ to $\omega_{32} + 2\pi \times 40\text{ MHz}$ in frequency, the angular frequency range where atoms can be rephased will be approximately $\Delta \in [-30\times 2\pi \text{ MHz}, 30\times 2\pi \text{ MHz}]$. Longer control pulses (of length $\sim 1\text{\mu s}$) with a slightly greater chirp range can rephase a range of $\Delta \in [-45 \times 2\pi \text{ MHz}, 45 \times 2\pi \text{ MHz}]$ without exciting unwanted atomic transitions. The two oscillator strengths are also not very

![FIG. 9. (Color online) Hyperfine energy-level spacings of $^{153}\text{Eu}^{3+}:\text{Y}_2\text{SiO}_5$ for the $^{153}\text{Eu}$ isotope (from [44]) on the left and the $^{151}\text{Eu}$ isotope (from [45]) on the right. The hyperfine levels used for the $\Lambda$ system (sketched in the middle) are connected to the relevant states with the broken blue arrows. Solid red arrows mark the optical transitions used.](image)
different ($D \simeq 2$), so driving both transitions in the adiabatic regime with the same pulse is feasible. With an optical lifetime of $T_1 = 2$ ms and spin coherence times exceeding days for temperatures around 2 K [46], this system seems fit for the realization of ROSE combined with spin-wave storage.

Of course, generating control pulses with a chirp spanning this range can be quite challenging. A possible alternative is to use the $^{151}$Eu isotope, where the hyperfine level spacing is about three times smaller [45]; see the right side of Fig. 9. Then $\omega_R = 2\pi \times 34.5$ MHz, and all resonance frequencies are higher than the $\omega_{12}$ frequency that belongs to the $|\pm 3/2g\rangle \leftrightarrow |\pm 5/2e\rangle$ transition except for $\omega_{32}$. We have $\omega_R = 2\pi \times 34.5$ MHz and the nearest unwanted resonance frequency is $\omega_{12} + 2\pi \times 46.2$ MHz belonging to the $|\pm 5/2g\rangle \leftrightarrow |\pm 5/2e\rangle$ transition. This system thus yields a somewhat smaller frequency range that can be used for signal storage, but requires a much smaller chirp range.

V. SUMMARY

In this paper, we have investigated coherence rephasing in an inhomogeneously broadened ensemble of $\Lambda$ atoms with three consecutive frequency-chirped control pulses for optical quantum memory applications. We have shown that, if the transform limited bandwidth is much smaller than the frequency difference of the lower energy levels ($\tau_p^{-1} \ll \omega_R$), but the overall bandwidth is greater, the control pulses can drive a cyclic permutation of the atomic populations in the adiabatic regime. With three such pulses interacting with the ensemble one after the other, it is possible to rephase the optical coherences left behind by a weak signal pulse, leading to the emission of a signal echo. We have shown that this rephasing, which happens when the atoms are predominantly in the ground state (i.e., the medium is not inverted), is analogous to the secondary rephasing of the ROSE scheme when two-level atoms are being rephased with two control pulses. There may also be a partial rephasing after one or two control pulses, when the medium is still inverted. Echo emission at this time can be prevented by choosing the control pulse propagation directions such that the primary echo fails the spatial phase-matching condition. At one point during this three-pulse rephasing process (after one or two control pulses, depending on the sign of $\omega_R$ and the chirp direction), the coherences left by the signal are stored as spin coherences between the two lower levels of the $\Lambda$ system. Thus this scheme of coherence rephasing conveniently incorporates long time spin-wave storage into the ROSE scheme using control pulses from a single source. The separation of the lower levels $\omega_R$ limits only the control pulse duration, but not the signal bandwidth that can be stored in the ensemble.

We have also investigated whether the current scheme is able to rephase the coherences of an optically dense storage medium. We have found, that despite the fact that the control pulses are distorted during propagation, coherence rephasing works well in a considerable domain in terms if spectral width and optical depth. The necessary condition for this is that the ensemble spectral width must be narrower than the bandwidth where the control pulses can rephase the atoms, i.e., the control pulses must cross both optical transitions for the entire spectral range of the ensemble.

We have also considered the application of the scheme in “naturally” inhomogeneously broadened media where the broadening is greater than the control pulse spectrum. In this case, however, there will be an $\sim \omega_R$ wide part of the ensemble where the control pulses interact with only one of the atomic transitions. On the one hand, this part of the ensemble will remain in the excited state after the third control pulse and will be a source of noise during signal retrieval. On the other, it will distort propagating control pulses much more in an optically dense medium, so the rephasing ability of the three pulses deteriorates fast. Thus the ensemble must be tailored to a width narrower than the control pulse spectrum before signal absorption. However, this does not mean that optical depth is lost at the signal frequency as in numerous other schemes.

Finally, we have argued that, to realize the scheme, we need materials whose excited-state sublevel separation is greater than the ground-state one and have shown that Eu$^{3+}$ doped into Y$_2$SiO$_5$ is a good candidate.