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Light transport in cold atoms and thermal decoherence

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By using the coherent backscattering interference effect, we investigate experimentally and theoretically how coherent transport of light inside a cold atomic vapour is affected by the residual motion of atomic scatterers. As the temperature of the atomic cloud increases, the interference contrast dramatically decreases emphasizing the role of motion-induced decoherence for resonant scatterers even in the sub-Doppler regime of temperature. We derive analytical expressions for the corresponding coherence time.

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Originally motivated by astrophysical purposes, wave transport in opaque media was first analyzed through a detailed balance of energy transfers within the scattering medium, leading to radiative transfer equations [1]. One main physical ingredient of this theory is a random-phase assumption, thereby discarding all possible interference effects. In its simplest form, this theory predicts a diffusive transport with spatial diffusion constant $D = \ell^2/3\tau$, where $\ell$ is the scattering mean free path and $\tau$ the transport time. However, it is now clear that interference can survive disorder average in the elastic regime and dramatically alter wave transport, leading for example to the physics of weak and strong localization [2]. There has thus been much effort in solid state physics to study, master and circumvent any possible phase-breaking mechanism to fully access the regime of coherent transport [3]. Coherence loss phenomena can be characterized by a phase-breaking time $\tau_\Phi$ and a corresponding coherence length $L_\Phi = \sqrt{D\tau_\Phi}$ beyond which interference effects are essentially washed out. The coherent transport regime (also known as the mesoscopic regime) is then reached when the medium size $L$ is smaller than $L_\Phi$. For instance, in metals or semi-conductors, it prevails up to $L_\Phi \approx 1 - 10\mu m$ even at very low temperatures and very pure samples, thus requiring the use of devices with micrometer size [3].

During the last two decades, the field got a renewed attention in systems using electromagnetic waves [4] and, among the large variety of scattering media employed so far, cold atomic vapors have recently emerged. An important feature of these media is the sharp resonance of the scattering cross-section (of order $\lambda^2$ on resonance, $\lambda$ being the wavelength), allowing for a continuous tuning of the scattering mean free path $\ell$ via the light frequency. Phase-breaking mechanisms in cold atomic vapors can be efficiently probed by using the coherent backscattering effect (CBS), a paradigmatic two-wave interference effect in multiple scattering [5]. In the limit of low light intensity, an important mechanism has been identified [6]. It is rooted in the Zeeman degeneracies of the atomic internal structure but could be healed by applying a magnetic field to the sample [7]. Another important phase-breaking mechanism is the residual thermal motion of atomic scatterers inside the cold vapor and is the main topic of this paper. Indeed, because of the Doppler and recoil effects, light frequency is changed upon scattering and transport is no more elastic. In previous papers [5,8], we have shown how the incoherent transport is affected by such effects. For typical alkali magneto-optical traps (MOT), Doppler-induced frequency shift is the dominant effect. Not surprisingly, an important parameter is the ratio of the typical Doppler shift $kv$ (being the 1D rms velocity, and $k = 2\pi/\lambda$ the incoming light wavevector) to the width $\Gamma$ of the atomic resonance. When $kv \gg \Gamma$, a single scattering event is enough to bring the photon completely out of resonance thereby altering transport significantly [6]. When $kv \ll \Gamma$, a single scattering event only slightly modifies the photon frequency and leads, as a cumulative effect, to diffusion in frequency space [3].

In this Letter, we report the first unambiguous experimental evidence of a phase-breaking mechanism in light transport induced by the residual atomic motion [1]. We first describe the experimental procedure and our main result. We then give an analysis of the physical ingredients at the heart of the light coherence reduction. We finally compare our data to the Monte-Carlo simulations described in [12], appropriately modified to include the effect of the atomic velocity distribution [3].

Our CBS experimental setup has been described in details elsewhere [12], the main difference here being the heating procedure to increase $v$. It is achieved by switching off the magnetic gradient of the MOT and then exposing the cloud to a $200\mu s$ long, slightly red-detuned, optical molasses. An increasing amount of heating is obtained by tuning the molasse frequency closer to resonance. The resulting 3D velocity distribution is measured by releasing atoms from the trap and imaging their ballis-
Monte-Carlo calculation assuming either a Gaussian (dashed line) or a Lorentzian (solid line) atomic velocity distribution. The CBS peak is recorded in the helicity non-preserving channel, with resonant laser light ($\delta = 0$). The optical thickness of the cloud is set constant at $b \approx 13$. For Rubidium, the velocity scale $\Gamma/k$ is 4.6 m/s.

![FIG. 1: CBS enhancement factor (full circles) as a function of the 1D typical atomic velocity $v$ and its comparison to a Monte-Carlo calculation assuming either a Gaussian (dashed line) or a Lorentzian (solid line) atomic velocity distribution. The CBS peak is recorded in the helicity non-preserving channel, with resonant laser light ($\delta = 0$). The optical thickness of the cloud is set constant at $b \approx 13$. For Rubidium, the velocity scale $\Gamma/k$ is 4.6 m/s.](image)

FIG. 1: CBS enhancement factor (full circles) as a function of the 1D typical atomic velocity $v$ and its comparison to a Monte-Carlo calculation assuming either a Gaussian (dashed line) or a Lorentzian (solid line) atomic velocity distribution. The CBS peak is recorded in the helicity non-preserving channel, with resonant laser light ($\delta = 0$). The optical thickness of the cloud is set constant at $b \approx 13$. For Rubidium, the velocity scale $\Gamma/k$ is 4.6 m/s.

![FIG. 2: (color online) CBS dephasing induced by the atomic motion, in the simplest case of two atoms moving with different velocities $v_1$ and $v_2$. The CBS effect builds on the two-wave interference between amplitudes associated to reversed multiple scattering paths (arrows). The Doppler effect induces frequency redistribution of the scattered light. Although the outgoing frequencies are identical, the intermediate frequencies differ, yielding a phase coherence loss and thus a reduction of the CBS contrast. A complex amplitude both incorporating scattering and propagation in the effective medium. For scatterers at rest, with no Zeeman degeneracies, these amplitudes are exactly balanced at backscattering (outgoing wave vector $-k$) yielding full CBS contrast. For Zeeman degenerate atoms, like Rubidium, we recover the results of [1]. Let us now consider the impact of the atomic motion. Because of the Doppler effect, the light frequency in the laboratory reference frame is modified. For the direct path, it is $\omega_d = \omega + (k' - k).v_1$ where $k'$ is the intermediate wavevector. After scattering by atom 2, the outgoing frequency is $\omega' = \omega + (k' - k).v_1 - (k + k').v_2$. For the reverse path, the frequency between the two atoms is $\omega_r = \omega - (k' + k).v_2 \neq \omega_d$, while the final frequency is again $\omega'$. Thus, although the outgoing frequencies are identical and do interfere, the intermediate frequencies differ by an amount $\approx kv$, resulting in different complex amplitudes and reducing on average the CBS interference contrast. Similarly, the frequencies seen by the atoms in their rest frames are different along the direct and reverse paths, leading to different scattering phaseshifts. More specifically, for small velocities $kv \ll \Gamma$ one can estimate the typical phase difference $\Delta \Phi$ induced by the atomic motion between the reversed paths:

$$\Delta \Phi = \Delta \Phi_s + \Delta \Phi_p \approx [\partial_x \Phi_s + k\ell \partial_x n]kv \approx kv/\Gamma$$

The first term corresponds to the scattering phaseshift, while the second describes the phaseshift associated to propagation over a distance $\ell$ in the effective medium. Although approximate, eq. (1) provides useful insights about the impact of $\delta$ at finite $v$ (see below). For larger scattering orders ($N > 2$), the elementary scattering and propagation processes have to be chained and randomly distributed phase differences will accumulate, decreasing...](image)
further the CBS contrast, see Fig. 3. In order to accurately compute the shape and height of the CBS peak, one must average over all atomic degrees of freedom, both external (position and velocity) and internal (motional quantum number), but one must also incorporate the experimental geometry such as the shape, size, optical thickness of the cloud as well as the waist and linewidth of the CBS probe laser beam. A Monte-Carlo approach as described in [23] is then necessary, where we now include the effect of the atomic velocity distribution 3. As mentioned earlier, some of the measured velocity distributions exhibited clear deviations from a Gaussian with an excess of fast atoms. Such atoms are responsible for large Doppler effects and are thus likely to kill CBS more efficiently. We have used in our simulations either a purely Gaussian velocity distribution or a Lorentz-like distribution (where we define \( v = v_0/(\sqrt{8\pi n^2}) \), \( v_0 \) being the FWHM-velocity). The results, displayed in Fig. 4, show a good qualitative agreement, with a fast decay of the enhancement factor with increasing \( v \). However, the Gaussian distribution show deviations at low \( v \) while the Lorentz distribution gives a better agreement. This pinpoints the important role played by a small fraction of atoms with large velocities. A quantitative agreement would require a better knowledge and control of the initial atomic velocity distribution. As can be seen, the CBS contrast decreases faster than naively expected. This is because higher scattering orders contribute significantly to the CBS effect. Indeed, at large optical thickness (\( b > 10 \)), they contribute altogether to \( \approx 30\% \) of the CBS peak height. As increasing scattering orders are more and more sensitive to motion-induced phase-breaking, this explains the reported observations.

Since the CBS scrambling originates from the frequency-dependent response of the sample (which is maximum around resonance), one could expect to recover a full interference contrast by detuning the laser at \( |\delta| \gg k\nu \). This is an important issue for the prospective CBS observation using a room-temperature vapor. Fig. 3 displays Monte-Carlo simulations of the CBS contrast as a function of the rms velocity \( v \) at \( \delta = 0 \) and \( \delta \gg k\nu \). Calculations are made for \( N = 2 \) (squares) and \( N = 40 \) (circles) in a semi-infinite medium. As can be seen (curves a and b, \( N = 40 \)), at low velocities \( k\nu \ll \Gamma \) the interference contrast is nearly frequency independent. This behavior is due to a compensation between the variations of the scattering and propagation terms. Indeed, the phase difference in eq. (1) is essentially proportional to the transport time \( \tau \) (bracketed term), which was shown to be frequency-independent near the resonance and equal to \( \Gamma^{-1}[2] \). As mentioned earlier and expressed in eq. (2), the decay rate is faster for higher scattering orders. At larger velocity (curves 1 and 2, \( N = 2 \)), Fig. 3 shows an increased frequency dependence of the interference contrast, which we interpret as an unperfect balance between the scattering and propagation phase-shifting in eq. (2). The interference contrast is partially restored in the \( |\delta| \gg k\nu \) limit, where the motion-induced decoherence is now essentially driven by the propagation term in eq. (1) of order \( k\nu/\Gamma \). The interference thus still decreases towards zero with increasing \( v \).

Beyond the specific case of CBS, one can quantitatively characterize the motion-induced phase breaking by defining the phase-breaking time \( \tau_\delta \) and its associated coherence length \( L_\delta \). We discuss here the simplest low-velocity case, \( k\nu \ll \Gamma \). At each scattering event, the photon de-tuning is slightly modified and this cumulative process turns into a random walk at sufficiently large scattering orders \( N \gg 1 \). Its dispersion is \( \approx k\nu \sqrt{N} \) which we also assume to be \( \ll \Gamma \). If the atomic velocities are Gaussian distributed and independent, the averaging over the degrees of freedom can be performed analytically, and the resulting interference contrast decreases like:

\[
\epsilon_N(v) = \exp \left[ -\frac{N^3}{12} \left( \frac{k\nu}{\Gamma} \right)^2 \right]
\]

Note that this quantity is independent of the initial de-tuning \( \delta \) as discussed before. This expression is also valid for larger velocities, in the off-resonant limit \( \delta \gg \sqrt{N} k\nu \) where the convolution by the velocity distribution can be neglected. The decay of the interference contrast described by eq. (2) is not exponential with \( N \), in contrast to other sources of decoherence like the atomic internal structure [2]. Nevertheless, one can still define a critical scattering order \( N_c = 3(3k\nu/2\Gamma)^{-2/3} \). This means
that the motion-induced phase-breaking mechanism can be characterized by:

\[ \tau_{\Phi} = 3 \left( \frac{3 k v}{2 \Gamma} \right)^{-2/3} \Gamma^{-1} \]  
\[ L_{\Phi} = \left( \frac{3 k v}{2 \Gamma} \right)^{-1/3} \ell \]  

Therefore, the mesoscopic condition \( L_{\Phi} > L \) reads:

\[ \frac{k v}{\Gamma} < \frac{1}{\ell} \]  

For our lowest temperature of \( \approx 40 \mu K \) (\( v = 6.5 \text{ cm/s} \)), eq. (3b) yields \( L_{\Phi} \approx 4 \ell \) and condition (4) is fulfilled for optical thicknesses \( b \leq 4 \).

At this point, an interesting comparison with electronic transport in metals can be made. There, the temperature dependence of \( \tau_{\Phi} \) is governed by several different mechanisms: electron-electron scattering (\( \propto T^{-2/3} \)), electron-phonon scattering (\( \propto T^{-3} \)) and spin-flip scattering (weakly \( T \)-dependent) which dominates at low \( T \) [1, 13]. In our case, \( \tau_{\Phi} \) is set by the internal structure \( \Phi \) (independent of \( T \)) and by the motion-induced decoherence, which, according to eq. (5), yields \( \tau_{\Phi} \propto T^{-1/3} \). At \( T = 40 \mu K \), \( \tau_{\Phi} \) is mostly determined by the internal structure (see Fig. 1).

A simple dynamical picture of the motion-induced coherence loss has been given by Golubentsev [16]. In this picture, phase-coherence is lost when the length of a multiple scattering path varies (due to the motion of the scatterers) by \( \lambda \) during the time taken by the light to follow this path. Golubentsev found expressions equivalent to ours, with however a huge difference: for the non-resonant case he considered, the transport time is \( \tau = \ell/c < 1 \text{ ps} \) while for our narrow resonance this time is \( \Gamma^{-1} \approx 27 \text{ ns} \), more than 4 orders of magnitude higher.

It is only because photons are “slowed down” inside the resonant medium that we are able to observe the destruction of phase coherence.

In conclusion, we reported the first observation of a motion-induced phase breaking of the CBS effect. The observed CBS reduction is due to the fast-varying frequency response of the medium in the vicinity of a resonance, which enhances the impact of frequency redistribution caused by Doppler effect. The rapid decrease of the interference contrast indicates that the phase coherence length \( L_{\Phi} \) associated with the residual motion of atoms in our coldest sample is not very large compared to the scattering mean free path, and smaller than the actual size of the cloud. According to the mesoscopic criterion derived in this paper, extremely low temperatures, in the nK range, are probably required to fully access interference effects in optically-thick cold atomic gases.

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