Renormalization group analysis of near-field induced dephasing of optical spin waves in an atomic medium

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Abstract

While typical theories of atom–light interactions treat the atomic medium as being smooth, it is well-known that microscopic optical effects driven by atomic granularity, dipole–dipole interactions, and multiple scattering can lead to important effects. Recently, for example, it was experimentally observed that these ingredients can lead to a fundamental, density-dependent dephasing of optical spin waves in a disordered atomic medium. Here, we go beyond the short-time and dilute limits considered previously, to develop a comprehensive theory of dephasing dynamics for arbitrary times and atomic densities. In particular, we develop a novel, non-perturbative theory based on strong disorder renormalization group (RG), in order to quantitatively predict the dominant role that near-field optical interactions between nearby neighbors has in driving the dephasing process. This theory also enables one to capture the key features of the many-atom dephasing dynamics in terms of an effective single-atom model. These results should shed light on the limits imposed by near-field interactions on quantum optical phenomena in dense atomic media, and illustrate the promise of strong disorder RG as a method of dealing with complex microscopic optical phenomena in such systems.

1. Introduction

The interaction of light with atomic ensembles provides the basis for numerous potential applications, such as quantum memories for light [1, 2], quantum nonlinear optics with strong photon–photon interactions [3–5], and quantum metrology [6–11]. In order to avoid the complexity associated with the large microscopic number of degrees of freedom, such as the large atom number and their positions, our standard theories for such systems typically favor a macroscopic approach. For example, for atom–light interactions in free space, the Maxwell–Bloch equations (MBE) [12–16] treat the atoms as a smooth polarizable quantum medium. The MBE have yielded many important insights into the physics that enables the applications above, as well as elucidating performance limitations [12, 17, 18].

Beyond macroscopic phenomena, many microscopic optical effects driven by granularity, dipole–dipole interactions and multiple scattering have been predicted, such as modifications of refractive indices and scattering rates [19–24], subradiance [25, 26], and coherent back-scattering [27–29]. Besides being of foundational interest, such microscopic effects could also have practical consequences on applications. For example, recently it was experimentally shown in reference [30] that such effects lead to a fundamental inhomogeneous broadening of optical transitions in an ensemble. This manifests itself as an additional dephasing on top of spontaneous emission decay for optical spin waves, with a rate that is exponential at early times and is directly proportional to atomic density. It was argued that this initial dephasing arises from the strong near-field interaction of a small fraction of particularly close nearest neighbors, quantitatively reproducing the experimental results. Separately, though, one might wonder what governs the
section 4, we present detailed numerical simulations of the spin wave dynamics from dilute to high-density media, and then formulate a simple, approximate, non-perturbative theory to predict a universal inhomogeneous broadening function for a disordered medium. From here, we will perform microscopic simulations of spin-wave dynamics. In section 3 we describe the RG approach, which enables us to describe the spin-wave dynamics in a microscopic way, taking into account atomic positions, near-field interactions, and multiple scattering of light, and which serves as the basis for the theoretical description of photon-mediated dipole–dipole interactions, which accounts for atomic details of individual atoms and their interactions.

We conclude and provide an outlook in section 5. We consider a minimal model consisting of many identical two-level atoms at fixed, random positions, near-field interactions, and multiple scattering of light, and which serves as the basis for the microscopic simulations of spin-wave dynamics. In section 3 we describe the RG approach, which enables us to predict a universal inhomogeneous broadening function for a disordered medium. From here, we then formulate a simple, approximate, non-perturbative theory to predict a universal inhomogeneous broadening function for a disordered medium. We conclude and provide an outlook in section 5. We consider a minimal model consisting of many identical two-level atoms at fixed, random positions, near-field interactions, and multiple scattering of light, and which serves as the basis for the microscopic simulations of spin-wave dynamics. In section 3 we describe the RG approach, which enables us to predict a universal inhomogeneous broadening function for a disordered medium. From here, we then formulate a simple, approximate, non-perturbative theory to predict a universal inhomogeneous broadening function for a disordered medium. We conclude and provide an outlook in section 5.

2. Microscopic model of atom–light interaction dynamics

We consider a minimal model consisting of $N$ identical two-level atoms at fixed, random positions $\{r_i\}_{i=1,...,N}$ that are uniformly distributed within a spherical cloud. The ground and excited states $|g\rangle$ and $|e\rangle$ have an electric dipole transition characterized by resonance frequency $\omega_0 = ck_0$ and wavelength $\lambda_0 = 2\pi/k_0$, and a single-atom excited-state spontaneous emission rate given by $\Gamma_0$. We also define a...
dimensionless density in terms of the number of atoms per cubic wavelength \( \eta = \lambda_0^3 N/V \), where \( V = \frac{4}{3} \pi R^3 \) is the volume of the ensemble and \( R \) its radius.

Within the standard assumptions in quantum optics (dipole, rotating wave and Markov approximations), the effects of photon-mediated dipole–dipole interactions, multiple scattering, and wave interference in spontaneous emission are captured by an effective atomic Hamiltonian \([31, 42, 43]\):

\[
H = -\frac{3\pi \Gamma_0}{k_0} \sum_{j} \mathbf{d}^i \cdot \mathbf{G}_0(\mathbf{r}_j - \mathbf{r}_i, \omega_0) \cdot \mathbf{d} \sigma_g^i \sigma_{ge}^j.
\]

(1)

Physically, the photon-mediated interactions between atoms are described by the free-space Green’s tensor \( \mathbf{G}_0(\mathbf{r}, \omega_0) \), the fundamental solution of the wave equation \([44]\). The Green’s tensor characterizes how a photon emitted by an atom at \( \mathbf{r}_i \) via the action of the atomic lowering operator \( \sigma_g^i = |g\rangle \langle e| \), propagates to a second atom at \( \mathbf{r}_j \), travelling a distance \( r_{ij} = |\mathbf{r}_j| - |\mathbf{r}_i| \). Moreover, as \( \mathbf{d} \) here represents the orientation of the atomic dipole matrix element, which we fix to be along \( \mathbf{e} \), we conveniently define \( \theta_{ij} \) as its angle with respect to \( \mathbf{r}_{ij} \). Doing so, the Hamiltonian \( H = \sum_{\ell} H_{\ell} \sigma_{g_{\ell}} \sigma_{ge_{\ell}} \), is characterized by the coefficients

\[
H_{\ell} = -\frac{3\Gamma_0}{4} e^{\imath k_0 r_{ij}} \left[ \frac{3 \cos^2 \theta_{ij} - 1}{(k_0 r_{ij})^3} - \frac{3 \cos^2 \theta_{ij} - 1}{(k_0 r_{ij})^3} - \frac{\cos^2 \theta_{ij} - 1}{k_0 r_{ij}} \right].
\]

(2)

The Hamiltonian is symmetric and non-Hermitian as a consequence of \( \mathbf{G}_0 \) enforcing reciprocity and being a complex quantity, including both coherent and dissipative interactions.

Important to later discussions, the real part of the Green’s function (describing coherent interactions) contains a near-field component, which dominates at small inter-atomic distances \( (k_0 r < 1) \). This near-field interaction explicitly reads:

\[
H_{\ell}^{\text{near}} = -\frac{3\Gamma_0}{4(k_0 r_{ij})^3} \left( 3 \cos^2 \theta_{ij} - 1 \right).
\]

(3)

The dissipative part instead describes collective spontaneous emission as arising from wave interference of the emitted light, while in the limit of a single atom, predicts the known spontaneous emission rate \( \Gamma_0 = \omega_0 d_g^2/3\pi\hbar\varepsilon_0 c^3 \), with \( d_g \) being the amplitude of the dipole matrix element for the atomic transition.

Generally, the presence of dissipation requires a master equation treatment \([31, 45–47]\), but the non-Hermitian Hamiltonian (1) is sufficient to describe the single-excitation regime of interest in our work, which generally reduces to solving a set of classical coupled dipole equations of motion \([16, 19, 43, 44, 48–52]\).

We will specifically be interested in applying the Hamiltonian (2) above to investigate the dynamics of a single-excitation ‘timed Dicke state’ or spin wave, defined as

\[
|k\rangle = \frac{1}{\sqrt{N}} \sum_j e^{\imath k \cdot r_j} |e\rangle.
\]

(4)

These collective states with well-defined wavevector constitute a natural basis to describe light–matter excitations. For example, phase-matched spin waves with \(|k| = k_0 = \omega_0/c\) are naturally and easily excited by an incoming resonant short pulse. By reciprocity, it is well known that they also efficiently emit into a narrow, well-defined direction centered around \( \mathbf{k} \) \([53]\), with a collectively enhanced rate \([16, 49, 50, 52, 54–58]\), \( \Gamma_0^{\text{coll}} \sim k_0 / \Gamma_0 = 1 + \text{ODF}/4 \), which linearly scales with the average optical depth of the medium. This narrow emission occurs due to constructive interference of the emitting atoms along the \( \mathbf{k} \) direction, and forms the basis of collective enhancement at the heart of efficient atom–light interfaces \([12, 59]\) and the applications mentioned in the introduction. This behavior can be equally derived by microscopic theories \([58]\) or by the macroscopic MBE \([16, 54, 55]\).

It should be noted that phase-matched spin wave excitations undergo non-trivial macroscopic spatio-temporal propagation dynamics \([60, 61]\). This makes it challenging to quantify the magnitude and effects of microscopic dephasing, due to the difficulty in defining an ideal time-evolving reference state to compare to, if dephasing could hypothetically be eliminated. An elegant, robust solution to this problem was proposed in reference \([30]\), with experimental realization based on a series of time-domain spin-wave control techniques \([58, 62]\).

The experimental technique relies on sending a series of very short (sub-ns) pulses to deterministically shift the spin wavevector, before the collective emission has a significant effect. These short pulses act on an additional transition \(|g\rangle - |a\rangle\), as represented in figure 2(a). To understand the basic mechanism it is sufficient to consider a single atom, at position \( \mathbf{r} \), initially prepared in the state \(|g\rangle + e^{\imath k \cdot \mathbf{r}} |e\rangle\), where \(|k| = k_0 \) is the matched wavevector of a probe field used to initially excite the \(|g\rangle - |e\rangle\) transition. Then, a
state returns to interest, classical coupled dipole equations. From the time-dependent state, we can construct two quantities of interest: a first quantity, \( P_k(t) \), which describes the survival of the spin-wave order. Importantly, it was shown to be a measurable quantity in the experiments of reference [30]. The amount of time that the system spends in the mismatched state, directly reveals the magnitude of the order of the control pulses. In particular, one can map a phase-mismatched spin wave back to a matched one. The peak intensity emitted by this matched spin wave, and specifically the dependence on the atomic position. This means that on the Bloch sphere, the evolution of the state encloses a non-trivial solid angle and thus picks up a geometric phase that depend on the atomic position. This means that on the Bloch sphere, the evolution of the state encloses a non-trivial solid angle and thus picks up a geometric phase \( \pm 2k_c \cdot r \) (with the sign \( \pm \) depending on the order of the pulses). The procedure can also be performed multiple times \( n \). Although this phase imprinting is a single atom effect, when applied to an ensemble, it is equivalent to shifting the initial wavevector of the spin wave to \( k_p \rightarrow k_p \pm 2n k_c \), as represented in figure 2(c). The process can also be reversed by exchanging the order of the control pulses. In particular, one can map a phase-mismatched spin wave back to a matched one. The peak intensity emitted by this matched spin wave, and specifically the dependence on the amount of time that the system spends in the mismatched state, directly reveals the magnitude of microscopic dephasing, as detailed in reference [30].

The mismatched spin-wave excitations \( |k\rangle \), characterized by \( |k| \neq k_0 \), constitute elegant initial states to investigate microscopic dephasing dynamics, due to the absence (on average) of macroscopic spatio-temporal dynamics. For example, these states neither couple to light efficiently nor have any preferred emission or propagation direction. Due to the lack of any direction in which the emitted field (averaged over random configurations) interferes, the average initial spontaneous emission rate of this state reduces to the single-atom value \( \Gamma_{\text{coll}}^{\text{spin}} = \Gamma_0 [25] \). Since this state does not exhibit any background macroscopic dynamics, \( |k\rangle \) itself serves as a reference to compare against the actual time-evolved state.

Specifically, we will be interested in the time evolution under the dipole–dipole Hamiltonian (1) as given by \( |k(t)\rangle = e^{-iHt}|k\rangle \). Since the initial state contains only a single excitation, as argued above, the dynamics can be efficiently evaluated numerically, with the resulting equations of motion equivalent to classical coupled dipole equations. From the time-dependent state, we can construct two quantities of interest,

\[
P_k(t) = \langle k(t)|k(t)\rangle \quad \text{and} \quad O_k(t) = |\langle k|k(t)\rangle|^2.
\]

The first quantity \( P_k(t) \leq 1 \) monotonically decreases and gives the total remaining excited state population at any time \( t \), with the rest having been irreversibly lost due to (collective) spontaneous emission. The second quantity, \( O_k(t) \), on the other hand, quantifies the overlap with the initial spin wave, and thus describes the survival of the spin-wave order. Importantly, it was shown to be a measurable quantity in the experiments of reference [30].

While we will present a more systematic analysis in section 4, we provide a visual example of the physics encoded in \( P_k(t) \) and \( O_k(t) \) in figure 3. In particular, we simulate the dynamics of an initial spin wave \( |k\rangle \), with \( |k| = 6k_0 \), in a particular configuration \( \{x_j\} \) of a disordered gas of \( N = 10^4 \) atoms with density \( \eta = 10 \), and in a spherical volume of radius \( R/\lambda_0 \approx 6 \). The evolved state \( |k(t)\rangle \) under equation (1) is calculated for several specific times \( t \). In figure 3(a) we consider all the atoms contained within a slice \( \Delta y = R/35 \) of the center of the cloud, and plot their positions in the \( x-z \) plane. The colors represent the accumulated phase of...
Figure 3. (a) Snapshots of the time evolution of a mismatched spin wave. We initially prepare a mismatched spin wave $|k\rangle$ ($|k\rangle = 6.0k_0$) in a particular configuration $\{r_j\}$ of a disordered gas at density $\eta = 10$, $N = 10^5$ atoms and radius $R/\lambda_0 \sim 6$. The state is then left to evolve under the dipole–dipole Hamiltonian (1) and the projection over the initial state is computed, $\langle k | k(t) \rangle = \sum_j |f_j| e^{i\phi_j}$, extracting the time dependent amplitudes and phases in the single atom basis. To create the snapshots above we consider all the atoms contained in a slice of size $\Delta y = R/35$, plot their position along the $xz$ plane, and color them according to their accumulated phase in time evolution $\phi_j$, as a consequence of the interaction, for different times. The global effect of this dephasing is represented in (b) where we plot the global overlap $O_k = |\langle k | k(t) \rangle|^2$ (blue curve) and population $P_k = \langle k(t) | k(t) \rangle$ (purple) of the same mismatched spin-wave. Quantifying the deviation of the overlap with respect to the predicted decay $e^{-\Gamma_0 t}$ (red dotted line) is the main purpose of this work.

Figure 4. Average decay rate $\Gamma_k = -2 \text{Im} \langle k | H | k \rangle$ of a spin-wave with momentum $k = (0, 0, k)$ for (a) different densities $\eta$ of a spherically uniform atomic cloud of $N = 10^4$ atoms. In (b) we evaluate the same quantity but keeping the density fixed to $\eta = 500$ and varying the number of atoms and the radius of the cloud. All the curves are averaged over 200 realizations.

Each atom relative to its initial value $e^{ik \cdot r_j}$, with a strong dephasing evident at time $\Gamma_0 t = 1.0$. In figure 3(b), we then plot $P_k(t)$ (purple) and $O_k(t)$ (blue) as calculated for the entire ensemble, along with the single-atom spontaneous emission $e^{-\Gamma_0 t}$ for reference (dashed red). It can be seen that while the initial decay of the total excited population $P_k(t)$ occurs at a rate $\sim \Gamma_0$ (confirming the absence of collective enhancement) before slowing down, the spin wave survival ratio $O_k(t)$ decays significantly faster than $\Gamma_0$, due to the dephasing illustrated in figure 3(a).

To practically study dephasing dynamics of a mismatched excitation, it is important to realize that the condition for mismatching actually depends on system size. In a sufficiently small system $R \to 0$, intuitively, phase differences $e^{ik \cdot \delta r}$ between atoms become negligible and all spin waves begin to resemble the symmetric $|k = 0\rangle$ state. Such a state then has superradiant properties reminiscent of the Dicke limit [53, 63]. To check the degree of phase mismatch, we evaluate the instantaneous average decay rate, $\Gamma_k = -2 \text{Im} \langle k | H | k \rangle$, and check that it is sufficiently close to $\Gamma_0$. In figure 4, we plot the decay associated to a particular spin-wave $|k\rangle$, for different densities, atom number and radii. These decay rates generically show two peaks at $\pm k_0$, which are associated with modes that efficiently emit because of the phase-matching condition. It can be seen that these peaks become narrower as the system size is increased (either at fixed density or fixed atom number), while for spin wavevectors that lie sufficiently far from these peaks the spin waves have a decay rate that indeed approach $\Gamma_0$. We observe in figure 4 that our specific choice $|k\rangle = 6k_0$ fulfills our mismatching requirement $\Gamma_k \sim \Gamma_0$ for the ranges of system sizes and densities to be explored in this work.
3. A renormalization group approach

While the microscopic model (equation (1)) can be numerically solved for moderate atom number, its complexity scales directly with the number of atoms \( N \), and with the number of configurations needed to obtain disorder-averaged results. Furthermore, the exact numerics does not directly elucidate the underlying physics. Motivated by that, here we introduce a simple model, based on a strong-disorder RG approach, which clearly identifies the role that near-field interactions have on the dynamics, and which allows the effects of such interactions to be captured by a simple effective, single-atom theory.

In a previous work [30], we derived a theory to understand the short-time dephasing rate of a spin wave for dilute ensembles with densities \( \eta \lesssim 1 \). We then found that the dephasing is primarily attributable to a small fraction of atomic pairs separated by a distance much smaller than the optical wavelength, \( k_0 r < 1 \) (highlighted in figure 1(a)), which strongly interact via their near fields (equation (3)). The \( \sim 1/r^2 \) scaling of the near field implies (in three dimensions) that the presence of other atoms is just a weak perturbation on top of the strong pairwise interaction, such that the pair can be separately and approximately diagonalized.

In the single-excitation manifold, diagonalizing the near-field interaction (3) of a pair yields symmetric and anti-symmetric eigenstates, \( |\pm\rangle = (|eg\rangle \pm |ge\rangle)/\sqrt{2} \), which experience opposite frequency shifts \( \omega_{\pm} = \pm \frac{3}{4k_0}(1 - 3 \cos^2 \theta) \) relative to the bare atomic transition frequency.

The time evolution for the two-body problem can now be studied in terms of its normal modes. Concretely, in the single atom rotating frame, \( e^{-i\omega_0 t} \), an initially prepared two-body spin-wave \( |k(t)\rangle \), will evolve as \( |k(t)\rangle = e^{-i\omega_0 t} |e_k\rangle^2 + e^{i\omega_0 t} |g_k\rangle^2 \), having defined the projections \( |e_k\rangle = (|k\rangle \pm |\pm\rangle) \). Although the magnitude of \( k \) might be constrained in an experiment, we can take the conceptual limit where \( k \to \infty \), or infinite mismatch. This implies that the phase \( e^{ik\eta} \) of each excited atom is effectively random (being infinitely sensitive to the specific atomic position), which implies that the actual spin wave should have on average equal overlap with the \( \pm \) eigenstates, i.e. that \( |e_k\rangle^2 \to 1/2 \). The dynamics can therefore be equivalently modeled by replacing the two original atoms with two new atoms of new resonance frequencies \( \omega_{\pm} \), that now do not interact anymore through the near field, but evolve with their ‘free’ inhomogeneous phases \( e^{-i\omega_{\pm} t} \) (see figure 1(a)).

In reference [30], the short-time dynamics of a many-atom, dilute ensemble were approximated by identifying a small fraction of close-by atomic pairs that evolved with the phases \( e^{-i\omega_0 t} \) as argued above, while the remaining atoms were assumed to undergo no evolution. By taking the position dependence of the function \( \omega_{\pm} (r) \) and combining with the known distribution function of separations \( r \) of nearest neighbors in a random ensemble, an initial decay of the spin wave survival order was predicted and measured to be exponential, \( |O_k(\delta t)|^2 = e^{-T \delta t^2} \), with a density-dependent rate \( \Gamma_k/\Gamma_0 = 1 + \xi \eta \), where \( \xi = 1/6\pi \sqrt{3} \) for two-level atoms.

Now, in a dense ensemble (figure 1(b)) or at longer evolution times, the basic picture of replacing close-by atomic pairs with new effective atoms of renormalized frequencies does not change. However, a key realization is that after an atom has been renormalized, it can still see another atom close by with which it can strongly interact (again highlighted by red circles in figure 1(b)). This allows yet another renormalization step to take place, which now will involve the diagonalization of a pair of atoms with the effective frequencies previously obtained. Whereas only a small fraction of atoms dictates the initial decay in a dilute ensemble, here, we must specify a general procedure valid for any density, to repeatedly and hierarchically identify the single most strongly interacting pair (including the possibility that the pair contains already renormalized atoms) and replace them with two new effective atoms.

Starting from that overall intuition, we now formulate the general principles and assumptions underlying our proposed RG scheme, before presenting a more detailed description of its implementation. We also present the principles of the RG scheme in a manner that goes beyond the particular goal of interest here, involving the study of dephasing dynamics. Succinctly put, our RG scheme consists of three principles:

(a) The system dynamics is exactly described by the total Hamiltonian of equation (2). Although we have provided the matrix elements in the basis given by individually excited atoms \( |e_j\rangle \), formally we are free to choose any other basis. In our RG scheme, we thus choose to work in the basis defined by the eigenstates of just the near-field interaction \( H_{\text{near}} \), as given in equation (3). Note that this is simply a choice of basis, and in particular, makes no assumptions about the relative contributions of the near-versus far-fields to the system dynamics.

(b) We assume that the many-atom eigenvalue distribution of \( H_{\text{near}} \) can be approximately obtained by a series of diagonalizations over atomic pairs, through a procedure specified below. The pairwise diagonalization, which significantly reduces the complexity, is motivated by the fact that the near field between a close-by atomic pair can dominate over the near-field interactions between this pair and all
other atoms combined. While this is an approximation, we can numerically verify that the agreement between the approximate and exact eigenvalue distribution is nearly perfect (e.g. see figure 5 and the related discussion).

(c) We now arrive at the third principle, the only one that cannot be fully justified. Strictly speaking, the eigenstates of $H^{\text{near}}$ will be somewhat delocalized over a set of atoms, as dictated by our identification of strongly interacting pairs. However, we assume that in terms of their interaction with light (via the far-field dipole–dipole interactions and from any possible external fields), these eigenstates respond effectively as if they were perfectly localized point dipoles. In particular, under this assumption, the original system becomes optically equivalent to a set of point dipoles that only interacts via the far fields, and which has a resonance frequency distribution corresponding to the eigenvalue distribution from (3). While the degree of validity of this approximation is hard to analyze on formal grounds, a partial justification of why it might work well is given in reference [41].

With these principles in mind and having anticipated that RG can involve the renormalization of atoms that have already been renormalized in previous steps, here, we consider the more general case of two inhomogeneous atoms of general resonance frequencies of $\omega_i$ and $\omega_j$, interacting via the near field, as described by the two-body Hamiltonian in the single excitation sector,

$$H_{jj}^{2b} = \langle \omega_{jj} \rangle 1 + \left( \frac{\delta \omega_{jj}}{H_{jj}^{\text{near}}} \frac{H_{jj}^{\text{near}}}{H_{jj} - \delta \omega_{jj}} \right).$$

For convenience, we have defined $\langle \omega_{jj} \rangle = (\omega_j + \omega_i)/2$, $\delta \omega_{jj} = (\omega_j - \omega_i)/2$. To quantify the strength of the interaction, we define the ratio between the off-diagonal and the diagonal elements,

$$K_{jj} = L_{jj} |H_{jj}^{\text{near}}|/(|\delta \omega_{jj}| + 1),$$

where the matrix $L$ keeps the information of whether a pair of atoms has already been renormalized ($L_{jj} = 0$) or not ($L_{jj} = 1$). This prevents a pair of renormalized atoms from being renormalized between themselves multiple times (although each atom from the pair can be renormalized with other atoms). Intuitively, a large value of $K_{jj}$ (which requires $L_{jj} = 1$) means that the strength of the interaction is able to further split the original frequency difference $\delta \omega_{jj}$. Thus, the most strongly interacting pair is identified as that with the largest value of $K_{jj}$ (red circles in figure 1). Once identified, the full diagonalization of (6) gives two eigenvalues, $\omega_{\pm} = \langle \omega_{jj} \rangle \pm \sqrt{\delta \omega_{jj}^2 + (H_{jj}^{\text{near}})^2}$. The pair can be therefore replaced by an approximately equivalent one, made of two atoms with the new resonance frequencies $\omega_{\pm}$ and that do not interact anymore through the near field (setting $L_{jj} = 0$).

Repeated application of this algorithm, which constitutes the RG flow, continues until all near-field interactions have been removed ($L_{jj} = 0$ for all pairs) and the atoms have been assigned the new effective frequencies $\{\omega_i\}$. In particular, this does allow for pairs with small values of $K_{jj} < 1$ to be renormalized, and we do not impose a cutoff to the RG based on the value of $K_{jj}$. This formally implements the principles of our RG approximation described earlier, where our goal is to describe the system in terms of the (approximately calculated) eigenstates of the near field. The actual numerical implementation of RG thus exhibits a complexity of $\sim N^2$ steps of renormalizing atomic pairs. However, as we will see in the next paragraph, RG gives rise to a universal distribution $P(\omega)$ of new effective resonance frequencies that is independent of atom number for sufficiently large $N$. This distribution can thus be used for all future
calculations at no complexity cost, e.g. in cases where $N$ is too large to make the direct numerical implementation feasible.

When the RG scheme is applied to multiple realizations of randomly distributed ensembles of atoms, we can build up the probability distribution $P(\omega)$ of the effective frequencies, which we illustrate in figure 5. As the near-field interaction only depends on distance through the dimensionless parameter $(k_{0}r)^{-3}$, the distribution when rescaled by density, $P(\omega/\eta)$, should be a universal function for a sufficiently large number of atoms and sufficiently large geometry, where boundary effects are negligible. We can directly confirm this numerically in figure 5, where we plot $P(\omega/\eta)$ obtained from RG for various densities.

The high-frequency tails of $P(\omega)$ correspond to the most strongly interacting pairs, which are renormalized at the beginning of the flow. This perfectly matches the simpler theory presented in [30] for dilute atoms, based on the probability distribution of nearest neighbors, where it was found that asymptotically

$$P(\omega) \sim \sqrt{\omega} \xi \frac{1}{2\pi\omega^2}.$$  \hspace{1cm} (7)

The central part of the distribution $P(\omega)$ instead consists of atoms that have been renormalized multiple times. In this sense, the presented RG scheme and the resulting probability distribution is the correct way to capture the near field induced inhomogeneous broadening of the medium and the induced dephasing rate of spin-waves in dense media, as we are going to discuss.

Formally, the RG procedure amounts to approximately diagonalizing the near-field part of the Hamiltonian (1) (an $N \times N$ matrix), by repeatedly identifying and diagonalizing a dominant interacting pair of atoms (a $2 \times 2$ block). We can quantify the error by comparing the resulting frequency distribution $P(\omega)$ obtained by RG, with the eigenvalue distribution obtained by exact, numerical diagonalization of the real, symmetric $N \times N$ Hamiltonian $H_{\text{near}}$ (equation (3)). It can be seen in figure 5 that the two are essentially indistinguishable, which validates principle (b) of the RG scheme.

As anticipated in principle (c), at the end of the RG flow, the resulting new effective (point-like) atoms can still interact through the far-field, which in general cannot be ignored or assumed to be a small effect. From here, one can solve the far-field dynamics in this new basis (which now no longer has near-field interactions). This was done for example in [41] to study the refractive index, finding excellent agreement between the optical response that can be calculated exactly from the full Hamiltonian of equation (2) and the one of an inhomogeneously broadened medium as obtained by the RG treatment.

In our particular problem of interest involving near-field induced dephasing, we essentially eliminate the effect of far-field interactions, due to the specific choice of a phase-mismatched spin wave as the initial state. We specifically recall that mismatched excitations are naturally decoupled (in an average sense) from free space, with no particular direction of emission and a decay rate $\Gamma_0$ free of collective enhancement. We will indeed see in the next section that far-field effects are essentially unnoticeable in the dephasing of mismatched spin waves while RG predicts the dephasing rate with great quantitative accuracy. Since decay can be treated separately now, RG results can be applied to predict the time evolution of a spin wave in an effective single-atom picture, focusing on the spin-wave survival ratio $O_{\text{sg}}(t)$. In particular, while the coherence of a single, isolated atom (without the rotating frame) is expected to evolve as 

$$\langle \sigma_{ge}(t) \rangle = \langle \sigma_{ge}(0) \rangle e^{-i(\omega_0 - \Gamma_0 t)/2},$$

the distribution in resonance frequencies of the new effective medium will result in an uncertainty of accumulated phase in time

$$O_{\text{sg}}(t) = \left| \int d\omega P(\omega) e^{i\omega t} \right|^2 e^{-\Gamma_0 t},$$ \hspace{1cm} (8)

thus introducing microscopically-driven dephasing due to inhomogeneous broadening. We emphasize that the simplicity of equation (8) arises from the elimination of macroscopic dynamics via our initial phase-mismatched state.

4. Spin-wave dephasing

We now present the exact numerical simulations and analysis of the time evolution of a mismatched spin wave for densities ranging from dilute ($\eta \ll 1$) to dense ($\eta \gg 1$), which we will then compare with the simple RG prediction of equation (8). To be concrete, we take an initial state consisting of a highly mismatched spin wave (equation (4)) with momentum $|k| = 6k_0$, $\hat{x}$-polarized and directed along $\hat{z}$, in an ensemble of $N = 10^4$ atoms. Then, the time evolution of the total excited-state population $P_k(t)$ and of the overlap with the initially prepared spin-wave order $O_{\text{sg}}(t)$ is calculated for $N_k = 500$ realizations of the disordered gas and averaged, as indicated by $\langle P_k \rangle$ for example. Numerical results are represented in figure 6,
where we also plot our prediction for time evolution of the overlap, $O_{\text{RG}}$, made in equation (8) (orange dash-dotted lines), based on the effective single-atom theory described in the previous section.

We first focus on the short-time dynamics. As introduced in the previous section, the short-time decay of a spin-wave is predicted to be exponential $[30]$, $\langle O_k(\delta t) \rangle \sim e^{-\Gamma_k \delta t}$, within a time interval $\Gamma_k \delta t \ll 1$. The rate $\Gamma_k = \Gamma_0 + \gamma$ is given by the sum of the single-atom emission rate, as predicted by the macroscopic MBE, and an additional density-dependent dephasing rate, $\gamma = \Gamma_0 \xi \eta$, with $\xi = 1/6\pi \sqrt{3}$. As shown in the inset of $\eta = 100$ (figure 6), for example, the microscopic dynamics reveal a short-time decay of the spin-wave order that is distinctly faster than the MBE prediction. The decay becomes even more evident at higher densities ($\eta = 10^5, 10^7$). We can confirm the density dependence in the short-time decay rate $\Gamma_k$ by fitting the curve $\langle O_k \rangle$ for each density in the short-time window, defined by $\Gamma_k \delta t < 0.1$, to an exponential of the form $e^{-\langle \Gamma_k \rangle \delta t}$, and plotting the dependence of the fit parameter $\gamma$ versus density parameter $\eta$ in figure 7(a). We also plot the prediction $\gamma = \Gamma_0 \xi \eta$ in red. It is important to observe that within our defined ‘short time’ interval, already $\sim 10\%$ of the initial spin-wave order is lost. An excellent agreement is observed over a large range of densities, both changing size and number of atoms (respectively blue squares and circles) of a random gas, confirming that this effect does only depend on the dimensionless density parameter $\eta$. This dephasing for moderate $\eta < 10$ was also observed experimentally in reference $[30]$.

At longer times and at higher densities, the decay of spin wave order noticeably deviates from exponential. Despite its simplicity, our single effective atom model based on RG (equation (8)), displays excellent agreement beyond the short-time interval. Viewed from the RG perspective, this non-exponential contribution comes from the frequency components near the center of the inhomogeneous broadening probability distribution $P(\omega/\eta)$ (figure 5), corresponding to atoms that are renormalized multiple times. Interestingly, at even longer times, it can be seen that for each density, the average $\langle O_k \rangle$ deviates from our prediction and saturates to a value that barely decreases over the range of times plotted. We furthermore observe numerically that this value closely coincides with the total excited state population remaining divided by the atom number, $P_k(t)/N$ (dashed purple curves). The slow decay of population $P_k(t)$ at long times is an effect that has been studied extensively in recent years, and is known as late-time subradiance $[26,32,30,61,64–66]$. While a microscopic derivation is difficult, a heuristic argument can be made that the remaining population should be roughly equally distributed throughout the ensemble, given a smooth initial distribution. Furthermore, given the randomness of the dynamics, this population will be
Figure 7. (a) Short-time dephasing rate vs dimensionless density $\eta$. Blue dots are extracted from an exponential fit at short times ($\langle O_k \rangle \sim e^{-\gamma_0 t}$), within the time interval $\Gamma_k \delta t = 0.1$, of the full time evolution of a mismatched spin-wave ($|k| = 6k_0$) at different densities $0.1 \leq \eta \leq 10^3$. The red line is instead the theoretical prediction for the short-time dephasing rate $\gamma = \Gamma_0 \xi \eta$, discussed in the main text. Simulations are performed with a cloud of atoms of fixed size ($R/\lambda_{eg} = 1.34$) and averaging over $N_s \sim 500$ realizations. (Inset) The same short time dephasing rate is evaluated (square dots), but now to explore the low-density regime we simulate the spin-wave dynamics in an ensemble with a fixed number of atoms, $N = 10^4$ (again averaged $N_s \sim 500$ times) and varying the radius of the cloud. (b) Numerical simulations of the time evolution of the averaged overlap $\langle O_k \rangle$ (solid blue lines) for a mismatched spin-wave ($|k| = 6.0k_0$) compared with the RG prediction $O_{RG}$ (orange dash-dotted line), and the total excited population divided by atom number, $\langle P_k \rangle / N$ (purple dashed line). The simulations consider a system of fixed density $\eta = 100$, but different atom number $N = 10^2, 10^3, 10^4$.

statistically evenly distributed among any $N$ extended modes that can be defined for the system, such as our spin wave mode of interest.

As far as we can numerically check (e.g. up to $N = 10^4$ atoms for a density of $\eta = 100$ in figure 7(b)), we see that the RG prediction $O_{RG}$ follows the actual spin wave survival ratio $O_k$ for increasingly long times as $N$ is increased, due to the decrease in the saturation value $P_k/N$. This strongly suggests that the single-atom RG prediction should be interpreted as the correct description of the dephasing dynamics in the thermodynamic limit, when the late-time population in any one mode $\sim 1/N$ becomes negligible.

5. Conclusions

In summary, we have developed an effective single-atom theory that describes well the non-exponential dephasing dynamics of optical spin waves in disordered atomic media, including at high densities and at long times. This theory is based upon the technique of strong disorder RG, which treats the potentially strong near field interactions in such a medium in a non-perturbative way.

We envision that our predictions, particularly in the high density regime, could be immediately explored using solid-state emitter ensembles such as rare earth doped crystals [67–70], where many atoms per cubic wavelength are typical. Separately, the remarkable accuracy by which RG is found to reproduce the dephasing dynamics suggests that it can be a powerful tool to quantitatively investigate and understand other microscopic optical phenomena in disordered systems.

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Data availability statement

No new data were created or analysed in this study.

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