Photon scattering from a cold, Gaussian atom cloud

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We study the effect of a weakly driven atomic cloud’s polarization distribution on its photon scattering line shape in the limit where the atoms can be treated as stationary and randomly distributed. We find three distinct polarization regimes. First, for dilute clouds, the polarization magnitude is relatively constant. Second, for denser clouds, polarization builds at the front of the cloud for near-resonant light. Third, when the cloud condenses to the point where its dimensions become comparable to the wavelength, light focuses towards the back of the cloud for red detuning. For these regimes, we show which “mean-field” frameworks accurately describe the differing photon-scattering line shapes. Finally, for even denser clouds, mean-field models become inaccurate and necessitate the full point dipole model that includes atom-atom correlations.

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I. INTRODUCTION

The ongoing study of the interaction of light and matter has been integral to understanding effects, such as superradiant spontaneous emission [1–4], coherent scattered radiation [2,5–15], collective Lamb shifts [16], and Anderson localization [17–19]. Because this research represents an abundance of different physical systems and phenomena, a diverse set of theoretical formulations is necessary to achieve an optimal understanding of each. The original work by Dicke [1], for example, showed that (for his idealized system) it is convenient to describe the cascade of radiation he termed superradiance with small symmetric subsets of the total Hilbert space, often termed Dicke states. In contrast, for weakly excited systems with more complex interactions, it is often advantageous to leverage the equivalence of the equations that govern a system of weakly driven atoms and coupled harmonic oscillators, Eq. (2) [5,14,20]. Here, termed the point dipole model, this framework has allowed theorists to explore systems containing over $10^4$ atoms without approximation [8].

Although the point dipole model can provide quantitative results for a large variety of systems, it is highly numerical and, thus, often provides insufficient intuition about the system in question. Also, the point dipole model can be computationally expensive, limiting the calculation to relatively few atoms. For both reasons, approximate methods are often used to understand the interaction of light with many atoms. One approximate technique that has allowed theorists to make predictions about clouds that contain too many atoms to handle numerically is to simulate a cloud that is denser but has less atoms than the actual experiment [10]. An example of an approximate “mean-field” technique that can lead to deep insights and analytic results assumes an evenly excited phase distribution throughout the system (timed-Dicke state) [2,6]. Mean-field approaches—that treat illuminated matter as continuous dielectrics [21]—are also particularly intuitive tools for understanding many of these same effects when they are applicable [22,23]. Because these ostensibly different mean-field techniques can provide unique information about a given system, it is important to understand which physical regimes they are applicable in and when the full point dipole model is necessary. There has been recent work towards understanding the relationships between these two approaches, for example, Refs. [24–26] showed that the traditional mean-field approaches fail to describe the resonance shift in a small slab of matter, and Ref. [27] explored the role of the disorder introduced by the point dipole model in coherently radiating clouds of atoms. The divergence of approximate mean-field theories, such as the timed-Dicke state and the dielectric model from the more accurate point dipole model—as well as the physics behind these divergences—remains largely unexplored, however.

The main purpose of this paper is to understand, both qualitatively and quantitatively, the interaction of a plane light wave with a Gaussian cloud of cold atoms in terms of the interrelation between the cloud’s polarization distribution and its photon-scattering line shape. The atoms are distributed with a cylindrically symmetric density, Eq. (1), which can describe both a spherically symmetric cloud and one that is elongated along the direction of laser momentum. We compare the continuous dielectric model with the full point dipole calculations to understand the processes that govern the results and to show the regime of validity of different approximations. We also show how photon scattering is modified by attenuation and focusing effects. In order to obtain results for a wide range of atom numbers, we implement a unique iterative numerical technique capable of simulating over $10^5$ atoms without approximation. We also implemented a paraxial solution of Maxwell’s equations of a plane wave interacting with...
a continuum dielectric with index of refraction proportional to a Gaussian.

The paper is organized as follows: Sec. II gives a qualitative description of the focusing and attenuation of light by the Gaussian dielectric. Sec. III contains the methods used in the calculations, Sec. IV contains the results of the calculations, Sec. V contains a short discussion of conclusions, and the Appendix contains information about the numerical method used to solve for the coupled dipoles, Appendix, Sec. 1, and a discussion of the paraxial approximation, Appendix, Sec. 2.

II. QUALITATIVE PROCESSES

Before presenting the detailed results, we describe the qualitative features that govern the attenuation and focusing of a plane light wave incident on a Gaussian dielectric.

We find three distinctive regimes of the atom cloud that depend on its density and shape and on the laser detuning (see Fig. 1), each regime characterized by its polarization distribution. The figure shows a continuum dielectric calculation of a plane light wave incident on a Gaussian dielectric, Sec. III contains the methods used in the calculations, Sec. IV contains the results of the calculations, Sec. V contains a short discussion of conclusions, and the Appendix contains information about the numerical method used to solve for the coupled dipoles, Appendix, Sec. 1, and a discussion of the paraxial approximation, Appendix, Sec. 2.

First, for clouds with small ODs, the atom excitations are nearly evenly distributed throughout the cloud for all laser detunings (see the OD = 1/4 case); thus, models that assume an even distribution [6] give good agreement with the full point dipole model. When the cloud becomes more dense, the laser intensity is substantially reduced as it traverses the cloud, causing excitation to be much more likely in the front of the cloud for near-resonant light, causing non-Lorentzian line shapes; this is exemplified by the OD = 4 case where Δ = 0, left column, has considerable attenuation whereas the detuned case, right column, has more uniform excitation. Neither case has substantial focusing. In this regime, the photon scattering is well described by a single parameter: OD. Finally, for clouds with dimensions comparable to a wavelength, the light focuses towards the back of the cloud for red detuning [8] (see the OD = 64 case), is defocused for blue detuning and is mainly absorbed for zero detuning. Because of focusing, the red detuned light has larger polarization for z > 0 through the center of the cloud whereas there is almost no polarization near the center of the cloud for zero detuning. For all of the cases in Fig. 1, the continuum dielectric model reproduces the total photon-scattering and the forward photon-scattering rates from point dipole calculations averaged over many spatial configurations, even the OD = 64 cases, although accuracy requires a solution of Maxwell’s equations that includes focusing.

FIG. 1. The continuum model calculation for the y = 0 cross section of the spatial dependence of |E|ρ, which is proportional to the polarization. In all plots, |E| has been divided by |E| at z → −∞, and the density is divided by the peak density; if there was no attenuation or focusing, then |E|ρ would have a maximum value of 1 at x = z = 0. In all calculations, the OD label refers to the optical depth for on-axis and on-resonance light. The spatial density is characterized by ξ = 2, Eq. (1), and N = 2.1. The detuning is set at Δ = 0 for the left column and Δ = −0.4(1 + OD/8)Γ for the right column to accentuate the focusing effect.
At even higher densities, we find differences between the continuum dielectric and the point dipole calculations; the Clausius-Mossotti equations do not improve the agreement between the point dipole and the continuum model and, in fact, tend to give worse agreement at higher densities as was found in Refs. [24–26]. This is due to the emergence of dipole-dipole correlations between atoms and diffraction perpendicular to the laser.

III. METHODS

This section describes the calculation of a plane wave of low intensity light interacting with a Gaussian cloud of atoms. This is performed using two separate formalisms. In Sec. III A, we describe treating the atoms as stationary and interacting through the point dipole Green’s function [8–12,24–28,30]. In Sec. III B, we treat the cloud as a continuum dielectric \( \chi_c \) with a Gaussian spatial dependence. For all calculations, we assume that the direction of propagation is \( z \) and the direction of polarization is \( x \): \( \vec{k} \parallel \hat{x} \) and \( \hat{\ell}_{\text{las}} = \hat{e}_z \). All of the calculations assume \( J = 0 \) to \( J = 1 \) transitions. Finally, this section assumes that the change in \( k \) over a resonance linewidth is much smaller than \( k \), which is an accurate approximation for optical transitions.

In our calculations, the atoms were given random positions following a Gaussian density distribution:

\[
\rho(x, y, z) = \frac{N}{(2\pi)^{3/2}r_f} e^{-[(x^2 + y^2 + z^2)/(2\xi^2)]},
\]

where \( N \) is the number of atoms, \( r_f \) is the geometric mean of the \( x, y, \) and \( z \) standard deviations of the atom cloud, \( \xi \) is a shape parameter, and the average density is \( \bar{\rho} = N/[(4\pi)^{3/2}r_f^3] \). \( \xi \) equals 1 for a spherical distribution and is greater than 1 for a cloud elongated in the direction of propagation. The cooperativity parameter \( b_0 = 3N/\bar{\rho} \) is related to the on-resonance OD through the center of the cloud: \( \text{OD} = \xi b_0 \).

A. Light scattering from atoms

In the weak-field limit, the effect of a monochromatic beam of wave-number \( \vec{k} \) and polarization \( \hat{\ell}_{\text{las}} \) on a cloud of atoms can be determined by the \( \langle a_j^{(a)} \rangle \equiv \langle \sigma_j^{(a)} \rangle \), i.e., the expectation value of the lowering operator for component \( j \) of the atom [5,14,20]. We assume that the atoms have the same thickness (the de Broglie wavelength of a typical atom is much smaller than the average separation between atoms) and are the same as the Doppler broadening of the transition is much smaller than the natural linewidth, and the typical distance traveled during a lifetime of the excited state is much smaller than the average separation between atoms). Under these conditions, the atoms can be approximated as randomly positioned but stationary:

The condition on the de Broglie wavelength means we do not have to treat the atom as spread through a large part of the cloud, the condition on the Doppler broadening means the scattered photons are not shifted out of resonance with other atoms, and the condition on the distance traveled means the strength of a dipole-dipole interaction does not strongly vary. In this limit, the equations of motion for the amplitude of oscillation are

\[
\frac{d a_j^{(a)}}{dt} = \left( i\Delta - \frac{\Gamma}{2} \right) a_j^{(a)} - \frac{i\Omega_R}{2} (\hat{\ell}_{\text{las}} \cdot \hat{e}_j) e^{i\vec{R}^{(a)}}
\]

\[
- \sum_{\alpha \neq \alpha} \sum_j g_{j,j'}(\vec{R}^{(a)},\alpha) a_j^{(\alpha)}
\]

where \( \alpha \) represents an atom index, the position of atom \( \alpha \) is \( \vec{R}^{(a)} \), \( \vec{R}^{(a)} = \vec{R}^{(a)} - \vec{R}^j \), and \( j, j' \) is indicating the component of the dipole from atom \( \alpha \) to atom \( \beta \). Here, \( \Omega_R \) is the transition Rabi frequency, \( \Delta \) is the detuning of the laser from the transition, and \( \Gamma \) is the decay rate of the excited state. \( \hat{e}_j \) is the unit vector in the \( j \) direction. The point dipole Green’s function \( g \) is given by

\[
g_{j,j'}(\vec{R}) = \Gamma \left[ \frac{\delta_{j,j'} h_0^{(1)}(s) + 3\vec{R} \cdot \vec{n}_{j'} h_2^{(1)}(s)}{2} \right],
\]

where \( h_0^{(1)}(s) = j_1(s) + in_1(s) \) are spherical Hankel functions of the first kind and \( s = \vec{k} \cdot \vec{R} [21] \). We solve for the steady-state \( \vec{a}^{(\alpha)} \) by setting the time dependence of Eq. (2) equal to 0 and solving the resulting matrix equation. When the number of atoms \( N \) was small, we numerically solved the linear equations using standard LAPACK programs that temporally scale \( \propto N^3 \). When \( N \) was larger than \( \sim 10^3 \), however, we solved for \( \vec{a} \) using an efficient \( \propto N^2 \) iterative method that we developed. This numerical technique enabled us to simulate clouds with more than 2 \( \times 10^4 \) atoms; the technique is described in the Appendix, Sec. 1.

The angular differential photon-scattering rate into \( \vec{k}_f \), normalized by \( \Omega_R^2/\Gamma^2 \), is given by

\[
\frac{d\gamma}{d\Omega} = \frac{\Gamma^2}{2\pi \Omega_R^2 N^2} \frac{[\vec{P}(\vec{k}_f)]^2 - [\vec{k}_f \cdot \vec{P}(\vec{k}_f)]^2}{[\vec{k}_f \cdot \vec{P}(\vec{k}_f)]^2},
\]

where \( \Omega \) is the solid angle and

\[
\vec{P}(\vec{k}_f) \equiv \sum_{\alpha} \vec{a}^{(\alpha)} e^{-i\vec{k}_f \cdot \vec{R}^{(\alpha)}}.
\]

Finally, the total scattering rate per atom, normalized by \( \Omega_R^2/\Gamma^2 \), is equal to

\[
\gamma = \int \frac{d\gamma}{d\Omega} d\Omega = -\frac{2\pi}{\Omega_R^2 N} \text{Re} \left[ \frac{\Omega_R}{2} \hat{\ell}_{\text{las}} \cdot \vec{P}(\vec{k}) \right],
\]

where \( \text{Re} \left[ \cdots \right] \) means to take the real component. The dimensionless form of the photon-scattering rate \( \gamma \) above is useful since the calculations are far from saturation and, thus, independent of the transition Rabi frequency \( \Omega_R \) up to a scaling factor.

B. Continuum model of photon scattering

To compare to the light scattering from stationary atoms, we solved Maxwell’s equations with a continuum electric susceptibility \( \chi_c \), giving the equations,

\[
\nabla^2 \vec{E} - \nabla (\nabla \cdot \vec{E}) + k^2 \vec{E} = -k^2 \chi_c \hat{x} (\hat{x} \cdot \vec{E}),
\]

where \( k \) is the wave number, \( \chi_c \) is the dielectric constant, and \( \hat{x} \) is the unit vector in the \( x \) direction.
where $\delta(k \cdot E)$ on the right-hand side accounts for the two-state approximation for the transition instead of all three components of $J = 1$.

For a $J = 0 \rightarrow 1$ transition, the low-density form of the electric susceptibility is [31]

$$\chi_e^{(d)}(\Delta) = \frac{\chi_e^{(d)}(0)}{1 - (2\Delta/\Gamma)} = \frac{i\rho \sigma / k}{1 - (2\Delta/\Gamma)}.$$  

(8)

where $\Delta$ is the laser detuning, $\sigma$ is the cross section for scattering photons out of the original direction, and $\rho$ is the density of atoms in Eq. (1). For a $J = 0 \rightarrow 1$ transition, the cross section is $\sigma = 6\pi/k^2$. The electric susceptibility for a perfect, homogeneous, and isotropic gas is given by the Clausius-Mossotti (or Lorentz-Lorenz) form [21]

$$\chi_e = \frac{1}{3} \chi_e^{(d)} \Rightarrow \chi_e(\Delta) = \frac{i\rho \sigma / k}{1 - (2\Delta/\Gamma)}.$$  

(9)

where $\Delta' = \Delta + (\Gamma \rho \sigma / 6k)$. This gives a density-dependent shift of the resonance of $-\pi \Gamma \rho / k^3$.

In the paraxial approximation [32], the scattered wave has a slow dependence in the direction transverse to $\vec{k}$. Assuming $\vec{k} = k\hat{e}_z$, and the polarization of the incoming light to be $\hat{e}_z$, the electric field is approximated by

$$\vec{E}(x, y, z) \approx \hat{e}_z e^{ikz} E_0 \psi_e(x, y, z),$$

(10)

where to lowest order,

$$i \frac{\partial \psi_e}{\partial z} = -\frac{1}{2k} \nabla^2 \psi_e - \frac{k}{2} \chi_e \psi_e,$$

(11)

with $\psi_e(x, y, z) \rightarrow -\infty = 1$, $\nabla^2 = \partial^2 / \partial x^2 + \partial^2 / \partial y^2$, and there is spatial dependence to $\chi_e$ from the density, Eq. (1). This equation arises from an expansion of Maxwell equations, Eq. (7), in powers of $f$ which is the ratio of scale size of the variation in $z$ (roughly $1/k$) to that in $x$ (roughly the width of the cloud); $f$ is explicitly defined below Eq. (A6) in Appendix Sec. 2. The derivation of this and higher-order terms are discussed in the Appendix, Sec. 2. Except for Fig. 9, the first- and second-order corrections did not significantly change the results, implying the paraxial approximation is accurate for the case discussed in Sec. IV C.

We also considered the eikonal approximation, which simplifies Eq. (11) as

$$i \frac{\partial \psi_e}{\partial z} = -\frac{k}{2} \chi_e \psi_e \Rightarrow \psi_e = \exp \left[ \frac{k}{2} \int_{-\infty}^{\infty} \chi_e(x, y, z') dz' \right],$$

(12)

which leads to an analytic $\psi_e$. As a result, the detuning dependence of the forward and total photon scattering is fully described by OD for systems where the eikonal approximation is accurate. This means that—in this regime—clouds with larger $N$ and smaller $\rho$ may be accurately described by clouds with smaller $N$ and larger $\rho$.

Using Eq. (10), the amplitude of oscillation for the $n$th atom is approximately,

$$\bar{a}^{(n)} = \hat{e}_z \frac{\Omega_R}{2\Delta + i\Gamma} e^{i\chi_e^{(n)} \psi_e(\vec{R})}.$$  

(13)

The sum in Eq. (5) is approximated as an integral,

$$\bar{P}(\vec{k}) \approx \frac{\Omega_R}{2\Delta + i\Gamma} \int \rho(\vec{r}) e^{i\delta(\vec{k} \cdot \vec{r})} \psi_e(\vec{r}) d^3r.$$  

(14)

This expression can be used in Eq. (4) to obtain the differential scattering rate in the forward direction or in Eq. (6) to obtain the total scattering rate. However, it cannot be used for angles substantially different from the forward direction because of the paraxial approximation and because it does not account for the random scattering from individual atoms, which dominates at larger angles.

The maximum $\chi_e$ is when $\Delta' = 0$ and $\bar{r} = 0$ which leads to $\chi_e \max = i\bar{b}^3/\sqrt{6\pi N}$. This can be used to estimate how well the low-density limit holds everywhere in the gas. However, this estimate can be misleading because the light does not reach the center of the cloud when $b_0$ is large and diffraction can be ignored. For light going on axis, the intensity is decreased by the factor of $\exp(-\xi b_0/2)$ when it reaches the center of the cloud for the spatially large clouds. However, atom clouds with smaller $N$ are also spatially small, and diffraction becomes increasingly important, see the OD $= 64$ ($b_0 = 32$) case of Fig. 1. We performed calculations for $b_0 = 8, 16, 24, 32$, and 40, although only results for 8 and 40 are given below. The $b_0 \geq 24$ cases have negligible intensity at the center of the cloud if diffraction effects can be ignored. However, for some of our parameters, diffraction is important, and light has non-negligible intensity at the cloud center for some of the large $b_0$ cases.

IV. RESULTS

We initially present results for the dilute gas limit in order to illustrate the effects of absorption and focusing by the cloud; these calculations require larger $N$ because $\chi_e \max$ is inversely proportional to $\sqrt{N}$ for a fixed $b_0$ and $\xi$, which necessitates our iterative numerical method (see Appendix, Sec. 1). We find that, in this dilute regime, the continuum model gives excellent agreement with the point dipole model. We then show that, for dense clouds, calculations based on classical electrodynamics treatments break down. This indicates the importance of correlations between neighboring atoms induced by dipole-dipole interactions.

A. $\xi = 1$, dilute gas limit

In this and the next section, all of the calculations of the paraxial approximation of the continuum use the low-density limit of the electric susceptibility, Eq. (8). This choice is explained in Sec. IV C.

The first results are for the total scattering per atom for different numbers of atoms for $b_0 = 8$ and 40 and $\xi = 1$; for the cases with $\xi = 1$, the OD $= b_0$. In Ref. [8], it was shown that the width of the resonance was $\Gamma' = (1 + b_0 \xi / 8)\Gamma$, so these calculations should give resonances from $2 \times 6\chi$ that of the single atom resonance when $\xi = 1$ and from $3 \times 11$ larger when $\xi = 2$. Calculations were also performed for $b_0 = 16, 24$, and 32 but are not reported because their properties can be inferred from the calculations described
agreement.

For $\xi = 1$ and for different cooperativity parameters $b_0$, calculated using Eq. (2). Each plot shows the results for two different numbers of atoms: solid (red) line $N = 2^{11}$ and long dashed (blue) line $N = 2^{17}$. For both plots, the different calculations are nearly the same so that the different lines are nearly indistinguishable. Plots for $b_0 = 16, 24,$ and 32 and for $N > 2^{11}$ showed a similar level of agreement.

Below. Results are reported for $2^{11} = 2048$ and $2^{17} = 131,072$ atoms. The calculations were averaged over many runs until a total of $2^{19}$ atoms were included in the calculation.

To give some rough sizes, the peak density times $\lambda^3$ gives the peak number inside a cubic wavelength. This is $(b_0^2 \pi / 3)^{3/2} / \sqrt{N}$. For $b_0 = 40$ and $N = 2^{17}$, there are $\sim 2$ atoms per cubic wavelength at the center of the cloud whereas $N = 2^{11}$ gives 17 atoms per cubic wavelength. If the more relevant quantity is the density times $1/k^3$, then, the maximum number is $(b_0/(6\pi \xi ))^{3/2} / \sqrt{N}$. By this quantity, all calculations in Fig. 2 have much less than one atom per $1/k^3$ ($\approx 0.07$ atoms for $b_0 = 40$ and $N = 2^{11}$).

Figure 2 shows plots of the total scattering rate per atom versus the detuning for two different cooperativity parameters: $b_0 = 8$ and 40. In each plot, there are two calculations for different values of $N$. The $N$ values are chosen to be $N = 2^{11}$ and $2^{17}$. Despite the drastically different parameters for each of the clouds, the overall line shape depends only on the value of OD. Although $N$ varies by a factor of 64, the total scattering rate per atom is essentially the same. This is because, for these parameters, the eikonal approximation, Eq. (12), gives very good agreement with the calculations from randomly placed atoms. There are two interesting trends to note. The first is that the line shape is changing with increasing $b_0$. For $b_0 = 8$, the line shape is approximated by a Lorentzian. However, for $b_0 = 40$, the central part of the line is narrower than for a Lorentzian. Also, the region near $\Delta = 0$ appears to have a dependence, such as $|\Delta|$, instead of $\Delta^2$, for $b_0 = 40$. Calculations were performed for $b_0 = 16, 24,$ and 32 with similar levels of agreement.

Figure 3 shows a comparison of three different calculations of the total photon-scattering rate $\gamma$ versus detuning for two different coherence parameters: $b_0 = 8$ and 40. The solid (red) line is using the full calculation Eq. (2) with $N = 2^{17}$ atoms in each run and was averaged until $2^{19}$ atoms were included. The long dashed (blue) line is from the paraxial approximation in Eq. (13) substituted into Eq. (14). The resulting $\tilde{P}(\tilde{k})$ was then used in Eq. (6). There were no adjustable parameters. The model calculation accurately reproduces the full calculation. The dashed (orange) line is a Lorentzian using the width $\Gamma' = (1 + b_0\xi/8)\Gamma$ with the height of the Lorentzian fit to
give agreement in the wings. Similar results were found for $b_0 = 16, 24,$ and $32.$

Figure 3 raises an interesting point. References [7,8,10] derived the broader linewidth Lorentzian (dashed orange line) by solving for the superradiant time dependence uniformly excited across the Gaussian distribution of atoms (timed-Dicke state). However, neither the paraxial nor the eikonal approximation of a continuum dielectric use this concept. The width in Eq. (13) is the single atom width $\Gamma$. The larger width emerging from Eq. (14) is solely due to the interplay of the nonuniform light intensity across the atom cloud as well as the phase change in Eq. (10). Figure 3 also shows that for larger values of $b_0$—when the polarization ceases to significantly penetrate the full cloud—point dipole and continuous dielectric models show a narrowing of the line-shape near-resonance; the timed-Dicke state models do not show this since the uniformly polarized state ansatz becomes insufficient.

The calculations allow us to untangle the coherent scattering of photons in the forward direction and the random scattering into large angles. To obtain the forward scattering rate, we integrated Eq. (4) over $\phi$ from 0 to $2\pi$ and integrated $\theta$ from 0 to $\cos(\theta_{\text{max}}) = 1 - 13.8/(k^2r_0^2)$. The $\theta_{\text{max}}$ was chosen so that forward scattering has decreased by, at least, two orders of magnitude from its maximum value. The result is shown in Fig. 4. There is a plateau in the forward scattering rate for a range of detuning around $\Delta = 0$. The atomic calculations of the forward scattering rate are, again, well reproduced by the paraxial approximation of the continuum distribution for all of the optical depths that were calculated. The eikonal approximation also agreed well with the atom calculation for the forward scattered photons except for at large $b_0$ and small $N$.

Figure 5 shows the full atom calculations as well as the paraxial approximation for the angular scattering rate per atom, Eq. (4) for $b_0 = 40$ and $N = 2^{17}$. This shows that coherently scattered light can be reproduced by the continuum dielectric model. The figure shows results from three different detunings. The $\Delta = 0$ case shows strong diffraction minima due to the strong scattering of light in the center of the cloud; these minima are too deep for the continuum calculation because it does not include the random scattering from pointlike atoms. At large detuning, the absorption is less, so the scattered light more closely follows a Gaussian form.

**B. $\xi = 2$, dilute gas limit**

When the Gaussian cloud is elongated in the laser propagation direction, there is more absorption and focusing of the laser beam. For $\xi = 2$ (i.e., OD $= 2b_0$), the total scattering versus detuning for $N = 2^{11}, 2^{13}, 2^{15},$ and $2^{17}$ are shown in Fig. 6. Unlike Fig. 2, the calculations with different numbers of atoms give different results for $b_0 \geq 24$ indicating the breakdown of the eikonal approximation. In all of the calculations, the scattering rates converge to a symmetric form as $N \to \infty$, but, for smaller $N$, the scattering rate is larger for $\Delta < 0$. In fact, there is a significant hump for $b_0 = 40$ at $\Delta < 0$ for the $N = 2^{11}$ calculation, the solid (red) line.
These new parametrical dependencies arise due to focusing of the light at $\Delta < 0$, which correlates with a breakdown of the eikonal approximation. The focusing bends the light skirting the edge of the cloud so that it interacts more strongly with atoms at the back of the cloud than would happen without focusing: Since more light goes through the cloud, there is more scattering. The focusing is also the cause of the more extreme case of the effect seen in Ref. [8] where atoms at the back of the cloud with $\xi \gg 1$ were more strongly excited than atoms at the front of the cloud. The effect is larger at smaller $N$ because the cloud is smaller and denser which leads to more focusing. Even more than $\xi = 1$, the large $b_0$ scattering rate has a dependence more similar to $|\Delta|$ than to $\Delta^2$.

These distributions are well reproduced by the continuum dielectric model that uses the paraxial approximation. Figure 7 shows the comparison between the atomic and the continuum model calculations of the total scattering rate for $N = 2^{11}$ and $2^{17}$; the continuum model calculations are nearly indistinguishable from the atomic calculations. Note that the hump at negative $\Delta$ for $N = 2^{11}$ is well reproduced.

The forward scattering rate for $\xi = 2$, $b_0 = 40$, and $N = 2^{17}$ and $2^{11}$ are shown in Fig. 8. This has a similar form to the spherical cloud, although the plateau starts at smaller $b_0$ (not shown). The results from the paraxial approximation to the continuum model are also shown. The $N = 2^{17}$ results are in excellent agreement, but there is a noticeable difference for $N = 2^{11}$ and small $|\Delta|$. The difference arises when the light diffracts back into the atom cloud. We found that including the next-order term did not improve the paraxial approximation which suggests that this difference is due to a breakdown of the continuum dielectric model for light propagation at higher density. The possibility that this difference arises because we use the low-density form of the susceptibility, instead of Eq. (9), is addressed in the next section.

C. Denser gases

Reference [24] showed that the Clausius-Mossotti (or Lorentz-Lorenz) form of the susceptibility, Eq. (9), does not
describe the model of light scattering from stationary atoms, Eq. (2). In all of the calculations above, the paraxial approximation of the continuum model used the low-density form for the susceptibility, Eq. (8). This did not make much difference in the calculations because the maximum of $\chi_0^{\text{id}}$ was not very large. Nevertheless, we also found that the calculations using $\chi_0^{\text{id}}$ were more accurate than using $\chi_0$ from Eq. (9) for smaller $N$.

Figure 9 shows a comparison between the atom calculation [the solid (red) line] and the paraxial equation using the low-density electric susceptibility [the long dashed (blue) line], Eq. (8), and the Clausius-Mossotti susceptibility [the dashed (orange) line], Eq. (9). The continuum calculation using the low-density electric susceptibility seems to overestimate the effect from focusing for $\Delta \sim -3\Gamma$ whereas the Clausius-Mossotti susceptibility is not accurate throughout the range $|\Delta| < 3\Gamma$. We found that the second-order correction to the paraxial approximation did not explain the difference for $N = 2^9$; although the correction to the paraxial approximation was not negligible for the $n = 2^7$ calculation, it could not explain the difference with the atom calculation. Overall, the low-density form of the susceptibility gives a more accurate representation of the total scattering versus detuning. The Clausius-Mossotti susceptibility gives a blue shift to the line whereas the low-density form gives a slight redshift due to focusing for $\Delta < 0$. The size of the effect is smaller than Ref. [24] because they used a much higher density and they used a uniform density whereas a Gaussian density was used in this paper. The calculations in Ref. [24] were for $\rho/k^3 = 2$ whereas Fig. 9 has $\rho_{\text{max}}/k^3 = 0.27$ for $N = 2^7$ and 0.14 for $N = 2^9$. These results show that the Clausius-Mossotti (or Lorentz-Lorenz) electric susceptibility does not reproduce the stationary atom calculation even in cases where $\rho/k^3 < 1$.

V. CONCLUSIONS

We have performed calculations of light scattering from a weakly driven Gaussian cloud of stationary atoms. We showed nontrivial effects on the scattering when moving from small to large optical depths. We also demonstrated a numerical method, Appendix, Sec. 1, that can solve for light scattering from many more atoms than is typical in current calculations. Thus, simulations can approach the number of atoms in experiments; results for up to $N = 2^{17}$ were presented.

We showed that the photon-scattering rate versus detuning is quite different from a Lorentzian at larger optical depths. This is because when polarization begins to build in the front of the cloud, the on-resonant forward scattered light does not propagate through and plateaus. For larger numbers of atoms, the total and forward scattering rates were quantitatively reproduced by a continuum model that used the low-density expression for the electric susceptibility. Even though it only contains linear absorption, the continuum dielectric calculation gave better agreement with the full point dipole calculations than models that use the single-photon superradiance framework. The full point dipole results for smaller atom number differ somewhat from the continuum model. Interestingly, worse results were obtained when using the Clausius-Mossotti (or Lorentz-Lorenz) form for the electric susceptibility, in agreement with the findings in Ref. [24].

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APPENDIX

Below are more detailed descriptions of the numerical methods used in the calculations above.

1. Iterative method

The steady-state solution of Eq. (2) involves the solution of a linear equation. For most of the calculations in this paper, we restricted the oscillators to only be in the $x$ direction which
means only the terms with \( j = x \) are included. The discussion in this section will focus on this case for simplicity, but it should be clear how to generalize to include all polarizations. For \( N \) atoms, this leads to an \( N \times N \) matrix equation of the form

\[
\sum_{b} A_{bb} \delta_{b} = s_{b}.
\]  

(A1)

For a small number of atoms (less than \( \sim 1000 \)), we used LAPACK subroutines to directly solve for \( a_{b} \). For a larger number of atoms, we used an iterative method based on successive over-relaxation.

The method proceeded in five stages. First, we ordered the atoms in the direction of laser propagation; for the \( \vec{k} = k\vec{e}_{z} \) used above, the atoms are ordered from smallest \( z \) to largest \( z \); the \( a_{b} \)'s are updated in this order so that the atom with most negative \( z \) is updated first and the atom with most positive \( z \) is updated last. Next, for each atom \( b \), we found the nearest \( M - 1 \) atoms \( b' \); these atoms will have the largest \( g \) (hence, the largest \( A_{bb} \)). The third step constructs an \( M \times M \) linear problem using the \( a_{b} \) from the previous iteration. The smaller linear system is defined by

\[
A_{bb} \delta_{b} = A_{bb} \delta_{b} \quad \text{for } b', b \in M,
\]

\[
\tilde{s}_{b} = s_{b} - \sum_{b \neq M} A_{bb} \delta_{b} \quad \text{for } b' \in M.
\]  

(A2)

We next solve the much smaller linear equation,

\[
\sum_{b} A_{bb} \delta_{b} = \tilde{s}_{b},
\]  

(A3)

using standard LAPACK subroutines and update only the atom \( b' \); \( a_{b} = (a_{b} + \tilde{a}_{b})/2 \).

We order the atoms from small to large \( z \) because atoms with smaller \( z \) affect those at larger \( z \) more strongly than vice versa. By taking them in order, the convergence speed was improved. More importantly, by directly solving Eq. (A3), we are able to account for the large coupling between close pairs (or triples or quadruples, etc.) of atoms.

We were able to converge all of the calculations with more than \( 2^{10} \) atoms using this method. Typically, we used nine iterations before convergence. Most of the calculations converged with \( M = 1 \) or \( 7 \) closest atoms. The calculations with the largest \( b_{0} \) and smallest \( N \) sometimes did not converge for \( M = 8 \) but did converge for \( M = 16 \). The calculation speed improves with smaller \( M \), so we first did all calculations with \( M = 8 \) and only repeated the failed ones with \( M = 16 \). The failed calculations were easy to determine because they had discontinuous jumps in scattering rate versus detuning.

This algorithm was much faster than directly solving Eq. (A1); direct solution is of order \( N^{3} \) operations whereas the iterative solution is of order \( N^{2} \) times the number of iterations. This also solved the problem of memory \( (A_{bb} \) has \( N^{2} \) complex numbers); when \( N \) was too large for the memory of our computer, we could compute \( A_{bb} \) on the fly instead of storing them in an array. Although not reported here, we performed calculations with \( N = 2^{18} \) atoms and one test calculation with \( N = 2^{19} \). The calculation with \( 2^{19} \) atoms would require an \( A \) with \( \sim 1/4 \times 10^{6} \) elements (i.e., over a terabyte of RAM) if performed by direct solution. Such large \( N \) can be reached because the density decreases with \( N \) which allows a smaller value of \( M \) to be used. The algorithm can be parallelized; most of the calculations were performed on a four-processor workstation, but the largest calculations were performed on a 20-processor workstation. No open source packages for implementing this algorithm currently exist.

### 2. Paraxial approximation

To derive the paraxial approximation as used in this paper, start from the expression for the exact Maxwell equation, Eq. (7), and substitute the form

\[
\vec{E} = E_{0}e^{i\vec{k} \cdot \vec{r}} \tilde{\psi} = E_{0}e^{i\vec{k} \cdot \vec{r}}(\psi_{T} + 2\psi_{C}),
\]  

(A4)

where \( \tilde{\psi}_{T} = a_{C} \tilde{x} + a_{S} \tilde{y} \) for any vector \( \tilde{a}_{T} \). This gives

\[
\frac{\partial \psi_{T}}{\partial z} + \frac{1}{2k} \vec{\nabla}_{T}^{2} \psi_{T} + \frac{k}{2} \vec{e}_{k} \cdot \vec{\nabla}_{T} \psi_{T} = -\frac{1}{2k} \vec{\nabla}_{T}^{2} \psi_{C} + \frac{1}{2k} \frac{\partial \psi_{C}}{\partial z} \psi_{T} + ik \psi_{T} + \frac{\partial \psi_{C}}{\partial z}.
\]  

(A5)

and

\[
\psi_{C} = \frac{i}{k} \vec{\nabla}_{T} \cdot \vec{\psi}_{T} - \frac{1}{k^{2}} \vec{\nabla}_{T}^{2} \psi_{C} + \frac{1}{k^{2}} \frac{\partial \psi_{C}}{\partial z} \vec{\nabla}_{T} \cdot \vec{\psi}_{T}.
\]  

(A6)

where \( \chi_{\vec{a}} \vec{\psi}_{T} = \chi_{\vec{a}} \psi_{T} \tilde{x} \). These equations are exact but are still difficult to solve. To simplify the equations below, we define the operator \( \mathcal{B} \) as

\[
\mathcal{B} \vec{\psi}_{T} = i \frac{\partial \vec{\psi}_{T}}{\partial z} + \frac{1}{2k} \vec{\nabla}_{T}^{2} \vec{\psi}_{T} + k \frac{1}{2} \chi_{\vec{a}} \vec{\psi}_{T}.
\]  

(A7)

We modify Eq. (A5) by substituting Eq. (A6) for \( \psi_{C} \) to give

\[
\frac{1}{k} \mathcal{B} \vec{\psi}_{T} = -\frac{1}{2k^{2}} \frac{\partial^{2} \vec{\psi}_{T}}{\partial z^{2}} + \frac{1}{2k^{2}} \vec{\nabla}_{T}^{2} \left( \frac{i}{k} \vec{\nabla}_{T} \cdot \vec{\psi}_{T} + i \frac{1}{k} \frac{\partial \vec{\psi}_{T}}{\partial z} \cdot \vec{\nabla}_{T} \frac{1}{2} \vec{\nabla}_{T} \psi_{C} \right).
\]  

(A8)

To obtain the paraxial approximation, one scales \( x, y \) by a width \( w \) and \( \psi \) by a length \( L = w/f_i \) (i.e., \( x = w \tilde{x}, y = w \tilde{y} \), and \( z = Lz \) with the barred coordinates being dimensionless). The ratio \( w/L = f_i \) is set equal to \( f \equiv 1/(k w) \). For the paraxial approximation, \( f_i \) should be small which means the distance scale of variations in \( x, y \) should be large compared to \( 1/k \), and the distance scale of variations in \( z \) should be large compared to that in \( x, y \). Substituting this scaling into the differential equations suggests that the three terms on the right-hand side of Eq. (A6) are of order \( f_i^3 \), \( f_i^2 \), and \( f_i \), respectively. The terms on the left-hand side of Eq. (A8) are of order \( f_i^3 \) or \( f_i^2 \), and, on the right-hand side, the terms are of order \( f_i \) if they involve \( \psi_{C} \) and \( f_i^4 \) if they involve \( \vec{\psi}_{T} \).

The functions are written as a series,

\[
\vec{\psi}_{T} = \vec{\psi}_{T}^{(0)} + \vec{\psi}_{T}^{(2)} + \vec{\psi}_{T}^{(4)} + \cdots,
\]

\[
\psi_{C} = \psi_{C}^{(0)} + \psi_{C}^{(2)} + \psi_{C}^{(4)} + \cdots.
\]  

(A9)

To obtain the equations for the different terms, one groups the same orders together. For example, since the first term on the right-hand side of Eq. (A6) is of order \( f_i^3 \), a term, such as \((i/k)\vec{\nabla}_{T} \cdot \vec{\psi}_{T}^{(4)} \) is of order 5 since \( \vec{\psi}_{T} \) is of order 4 and the
operation is of order 1. Equation (A6) is transformed to
\[
\psi^{(n)}_x = \frac{i}{k} \nabla_T \cdot \nabla_T^{(n-1)} \psi_T - \frac{1}{k^2} \nabla_T^2 \psi_T^{(n-2)} + \frac{1}{k^2} \frac{\partial}{\partial z} \nabla_T \cdot \nabla_T^{(n-3)},
\]
(A10)
when grouping terms of order \(n\). Defining the order of the \((1/k)^2\) operator is somewhat problematic due to the \(\chi_c\) term. We take it to be an order 2 operator, consistent with the two differential terms. Equation (A8) is transformed to
\[
\frac{1}{k} B \psi_T^{(n)} = -\frac{1}{2k^2} \frac{\partial^2 \psi_T^{(n-2)}}{\partial z^2} + \frac{1}{2k^2} \nabla_T \left(-\frac{i}{k} \nabla_T \psi_T^{(n-1)} + \frac{i}{k} \frac{\partial}{\partial z} \nabla_T \psi_T^{(n-2)} + \frac{\partial \psi_T^{(n-1)}}{\partial z}\right),
\]
(A11)
when grouping terms of order \(n + 2\). If \(n = 0, 1, 2\), the functions on the right-hand side can have negative superscripts. The rule for evaluating these are as follows: Any function with a negative superscript is zero everywhere.

The case discussed in the paper has the atoms only being polarizable in the \(x\) direction. This means all of the quantities of interest can be calculated from \(\psi_x\). Thus, through order 2, the equations to be solved are
\[
E \chi \psi_x^{(0)} = 0, \quad \psi_y^{(0)} = 0,
\]
\[
\psi_x^{(1)} = \frac{i}{k} \frac{\partial \psi_x^{(0)}}{\partial x},
\]
\[
E \chi \psi_x^{(2)} = -\frac{1}{2k} \left(\frac{\partial^2 \psi_x^{(0)}}{\partial z^2} + \frac{\partial [\chi_c \psi_x^{(0)}]}{\partial x^2}\right),
\]
(A12)
with \(\psi_x^{(0)}\) set to 0 as \(z \to -\infty\). Note \(\psi_x^{(2)}\) is nonzero, but it is not used in our calculations.


