Disordered ensembles of strongly coupled single-molecule plasmonic picocavities as nonlinear optical metamaterials

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We propose to use molecular picocavity ensembles as macroscopic coherent nonlinear optical devices enabled by nanoscale strong coupling. For a generic picocavity model that includes molecular and photonic disorder, we derive theoretical performance bounds for coherent cross-phase modulation signals using weak classical fields of different frequencies. We show that strong coupling of the picocavity vacuum with a specific vibronic sideband in the molecular emission spectrum results in a significant variation of the effective refractive index of the metamaterial relative to a molecule-free scenario, due to a vacuum-induced Autler-Townes effect. For a realistic molecular disorder model, we demonstrate that cross-phase modulation of optical fields as weak as 10 kW/cm² is feasible using dilute ensembles of molecular picocavities at room temperature, provided that the confined vacuum is not resonantly driven by the external probe field. Our work paves the way for the development of plasmonic metamaterials that exploit strong coupling for optical state preparation and quantum control.

I. INTRODUCTION

Strong light-matter coupling with single molecules in plasmonic picocavities has emerged as a resource for room-temperature quantum control with nanoscale optical fields. Organic chromophores in plasmonic picocavities [1–4] are promising platforms for studying cavity quantum electrodynamics (QED) at room temperature [5, 6]. Recent experiments [7] and rigorous theoretical modeling [8–11] have emphasized the quantum optical origin of commonly used plasmon-enhanced molecular spectroscopy techniques [12], offering new perspectives on conventional architectures that can stimulate the study of novel schemes for optical quantum control at the nanoscale [13].

Conventional molecular-cavity QED platforms based on planar optical microcavities exploit the interaction of the electromagnetic vacuum with an ensemble of vibronic coherences to reach the strong and ultrastrong coupling regimes [14–23]. The collective character of the interaction can enhance electric and charge transport processes [24–27], mediated by the cavity-induced delocalization of the molecular degrees of freedom involved in the transport process. Collective coupling can also lead to modifications of the chemical reactivity [28, 29] and optical response [30–33] of organic materials. Collective strong coupling in microway occurs through a mechanism analogous to dipole synchronization [34]. However, the local field that each individual molecule experiences in a microcavity is relatively small.

In contrast, in plasmonics, the extreme sub-wavelength field confinement achievable with current technology [35, 36] allows for light-matter interaction energies to overcome local thermal fluctuations, at the level of individual molecules. This can enable the implementation of local control protocols that exploit strong vacuum fields for studying optomechanical physics [7] and tailored photochemistry [37, 38]. Nanoparticle fabrication techniques can produce a large number of “molecular picocavities” [2]. The picocavity distribution can be strongly inhomogeneous, and must be sampled locally using tip-based nanopores [39–41], to extract spatially-resolved information about the light-matter coupling dynamics in the system. Although chemical methods are available to increase the homogeneity and reproducibility of the picocavity fabrication [4], the need to develop efficient local sampling method may be a challenge for the scalability and integrability of molecular picocavities in next-generation nanophotonic devices.

Instead of focusing on the local aspects of light-matter interaction in disordered picocavity ensembles, in Ref. 42 we explored a macroscopic approach in which cavity strong coupling was used for inducing nonlinear optical signals in the response of the ensemble. We assumed a scheme involving cis-trans molecular isomers (photoswitches) that are embedded in high-quality optical microwaves with photon lifetimes of several picoseconds. We exploited the unique spectral and coherence properties of molecular photoswitches to find suitable conditions for inducing a cavity-assisted transparency window in the absorption spectrum and implementing cross-phase modulation between external laser fields. The phase non-linearity was shown to be robust with respect to static disorder in the molecular dipole orientation and molecular transition frequencies, given a set of restrictions on the allowed vibrational and photonic dephasing times.

In this work, we significantly generalize the analysis in
Ref. [42] that could facilitate experimental implementations. We achieve this by reducing the number of physical assumptions imposed over the relevant molecular and photonic degrees of freedom, in particular, the type and properties of the cavity resonator structures needed for field confinement, and the class of organic molecules that couple to the vacuum cavity field. We now consider a broad class of organic chromophores that exhibit significant electron-vibration coupling in the lowest electronic singlet transition \(S_0 \leftrightarrow S_1\) for an intramolecular vinyl stretching mode of frequency \(\omega_v \approx 0.2\) eV [43]. We assume that static fluctuations of the molecular transition frequencies are the leading source of inhomogeneity in the ensemble, in an effort to understand the fundamental limits of coherent optical signals in a scenario where the energy disorder is dominant. Other types of inhomogeneity such as random dipole orientations have a smaller effect on the optical response of coupled light-matter systems [42].

On the photonic side, we adopt three simplifying assumptions about the properties of picocavities: (i) We consider a dilute ensemble of picocavity structures that are much smaller than the optical probe wavelength, and have negligible inter-particle interaction. This allows us to treat the ensemble as an effective medium which to lowest order is coherent optical signals in a scenario where the energy disorder is dominant. Other types of inhomogeneity such as random dipole orientations have a smaller effect on the optical response of coupled light-matter systems [42].

Building on these assumptions, we provide a proof-of-principle demonstration that single-molecule strong coupling with the picocavity vacuum induces phase modulation of a probe laser field that coherently drives a disordered ensemble with varying light-matter coupling strengths, inhomogeneous broadening of molecular transition frequencies, and sub-picosecond decoherence of molecular and photonic degrees of freedom. We also show that the presence of a second (signal) laser induces a controllable vacuum-enhanced cross-phase modulation, at intensities as low as a few kW/cm\(^2\). Such intensities are orders of magnitude lower than the light sources used in conventional nonlinear optics [44]. The ensemble nonlinearity can be optically gated and used as an optical switch, exploiting coherence at macroscopic scales despite the strong structural disorder.

In the rest of this article, we introduce the macroscopic approach that allows us to define the effective optical susceptibility of an ensemble of molecular picocavities (Sec. II A) and a nonlinear phase modulation signal (Sec. II B). Then we discuss the quantum electrodynamics model that describes the light-matter dynamics in individual picocavities (Sec. III), and finally study the dependence of the nonlinear phase modulation observables on the system parameters (Sec. IV V). We summarize our main results in Sec. VI.

II. EFFECTIVE MEDIUM APPROACH

The phase-changing plasmonic metamaterial studied in this work is illustrated in Fig. 1. We consider an inhomogeneous layer composed of an isotropic ensemble of plasmonic nanoparticle inclusions dispersed in a background thin film with dielectric constant \(\epsilon_d\). Such substrates are routinely used in plasmon-enhanced molecular spectroscopy [45–48]. In our case, we assume that the nanoparticles self-assemble as dimer picocavities with a single organic molecule embedded in the gap region, where strong light-matter interaction occurs locally within the electric-dipole approximation (see Fig. 2b). Spherical nanoparticle dimers are common [46], and single-molecule picocavities with a variety of geometries and material compositions can be produced [1–4].

We estimate the impact of the ensemble of single-molecule picocavities on the effective refractive index of the metamaterial. Due to the absorptive and dispersive character of molecular picocavities, the metamaterial index depends on frequency. In particular, we focus on the phase response of the system at the frequency of a weak probe field \(\omega_p\), which is resonant to the peak absorption frequency of the molecules, but is far-detuned from the mean gap resonance frequency of the picocavities \(\omega_c\). We consider molecules with large vibronic coupling such that the excited state reorganization energy is comparable with the intramolecular vibration frequency (\(\sim 0.2\) eV [49]). This ensures that molecules cannot strongly emit...
light at $\omega_p$, so the external probe laser is the relevant field source of the problem.

### A. Effective Index of the Metamaterial

Effective medium theories are used for evaluating the average macroscopic electromagnetic response of a spatially inhomogeneous medium [50]. For plasmonic nanoparticle ensembles, several techniques have been used for computing macroscopic dielectric response (effective index) [51], based on the well-known Maxwell Garnett theory [52, 53]. We follow an alternative effective medium methodology developed in Refs. [54, 55], which has been applied to describe inhomogeneous metals [56], polycrystal dielectrics [57] and polaritons in organic microparticles [58]. The dynamical variables of the problem are written as the sum of a macroscopic average and a local fluctuation, for which coupled equations of motion can be derived and solved.

We consider a dilute ensemble of $N$ independent single-molecule picocavities isotropically distributed over a background medium with dielectric constant $\epsilon_d$, as illustrated in Fig. 2a. The average picocavity size ($\sim 10$ nm) is much smaller than wavelength of the probe laser ($\sim 600 - 800$ nm), which is assumed to be the dominant electromagnetic source of the problem. In other words, inelastically scattered light from plasmonic nanoparticles as well as fluorescence and Raman scattering from organic molecules are negligibly weak at the probe frequency in comparison with the driving laser.

We start with the wave equation

$$\nabla^2 \mathbf{E}(\mathbf{r}, t) - \frac{1}{\epsilon_0 c^2} \frac{\partial^2}{\partial t^2} \mathbf{d}_d(\mathbf{r}, t) = \frac{1}{\epsilon_0 c^2} \frac{\partial^2}{\partial t^2} \mathbf{P}(\mathbf{r}, t), \tag{1}\nabla$$

where $\mathbf{D}_d(\mathbf{r}, t) = \epsilon_0 \mathbf{d}_d \mathbf{E}(\mathbf{r}, t)$ is the displacement field due to the background dielectric, and $\mathbf{P}(\mathbf{r}, t)$ is the polarization density due to picocavities with embedded dipoles, assumed to be large near the picocavities and vanishing in between. Computing this quantity from first principles is demanding, as it contains coupled charge density contributions from both the nanostructure and the embedded molecules [59]. Field enhancement factors due to interfaces can also in principle be obtained by solving Eq. (1) numerically given a distribution of particle geometries and compositions. However, we focus our analysis on the generalities of the coupled light-matter response at a particular frequency (probe) and defer the exact evaluation of the spatially-dependent local fields for future work.

We write Eq. (1) in the Fourier domain by expanding the electric field and polarization into components at discrete frequencies $\omega_n$, to read

$$\nabla^2 \mathbf{E}(\mathbf{r}, \omega_n) + \frac{\epsilon_d \omega_n^2}{\epsilon^2} \mathbf{E}(\mathbf{r}, \omega_n) = -\frac{\omega_n^2}{\epsilon_0 c^2} \mathbf{P}(\mathbf{r}, \omega_n). \tag{2}\nabla$$

For a weak probe field, the polarization at $\omega_n = \omega_p$ is expanded up to linear terms in the probe field, $E(\mathbf{r}) = E_p(\mathbf{r})$, as

$$\mathbf{P}(\mathbf{r}, \omega_p) = \epsilon_0 \chi(\omega_p, \omega_c, \omega_s, \mathbf{E}_c(\mathbf{r}), \mathbf{E}_s(\mathbf{r})) \cdot \mathbf{E}_p(\mathbf{r}, \omega_p), \tag{3}\nabla$$

where the susceptibility $\chi \equiv \chi(\omega, \mathbf{r})$ encodes the response at $\omega_p$ of the picocavities containing molecular dipoles.

In general, the susceptibility in Eq. (3) depends on the picocavity frequency $\omega_c$ and the local cavity fields strength, $\mathbf{E}_c(\mathbf{r})$. We also anticipate a dependence of the effective susceptibility on the frequency $\omega_s$ and amplitude $\mathbf{E}_s(\mathbf{r})$ of a signal laser field that is introduced in Sec. IV as an additional optical knob in the problem. In Sec. IV we derive explicit expressions for $\chi$ using a
quantum mechanical model for light-matter coupling in a picocavity.

Following Refs. [54–58], we write the probe field and susceptibility as \( (\omega \approx \omega_p \text{ is always assumed}) \)

\[
E_p(r) = \langle E_p(r) \rangle + \delta E_p(r), \tag{4} \\
\chi(r) = \langle \chi \rangle + \delta \chi(r). \tag{5}
\]

where \( \langle E_p(r) \rangle \) is the average probe field that propagates according to the effective index \( n(\omega_p) \), and \( \delta E_p(r) \) is a local field fluctuation, satisfying \( \langle \delta E_p(r) \rangle = 0 \). We expect that \( \delta E_p(r) \) are non-zero only in the immediate vicinity of the picocavities.

Similarly, the susceptibility is partitioned as a sum of a uniform average, \( \langle \chi \rangle \), and a local fluctuation \( \delta \chi(r) \), with \( \langle \delta \chi(r) \rangle = 0 \). Inserting Eqs. (3), (4), and (5) in Eq. (1), we can derive coupled equations for the field and susceptibility averages and their fluctuations. We solve these equations in Appendix A using perturbation theory, with the small parameter being the ratio of the typical inclusion scale to the wavelength of the electromagnetic wave in the medium. We find that the effective homogeneous index of the metamaterial at the probe frequency splits into three terms,

\[ n^2(\omega_p) = \epsilon_d + \langle \chi(\omega_p) \rangle + \Delta \epsilon_p, \tag{6} \]

where \( \langle \chi \rangle = (N/V)\chi_0 \), with \( \chi_0 \) being the susceptibility of a single inclusion, is the average susceptibility, and

\[ \Delta \epsilon_p(\omega_p) = \frac{\omega_p^2}{c^2 \sigma_d^2} \int \frac{dq}{2\pi} \frac{K(|k-q|)}{\epsilon_d + \langle \chi \rangle - q^2}. \tag{7} \]

is a correction to the homogeneous index, which takes into account spatial correlations between local fluctuations of the electric field and polarization, as captured by the Fourier transform of the spatial correlator \( K(|r_1-r_2|) = \delta \chi(r_1)\delta \chi(r_2) \).

Not going into detailed analysis of \( \Delta \epsilon_p(\omega_p) \), which can be done explicitly for specific experimental configurations of metamaterial, in Appendix A we argue that the correlator \( K(r) \) must vanish at the scales of the order of the inclusion size \( r_0 \sim 10 \text{ nm} \); this simply reflects the assumption that the picocavities are independent. Assuming \( K(r) \propto \exp[-r^2/(2r_0^2)] \) yields \( K(q) \propto \exp[-q^2r_0^2/2] \), which is vanishingly small as long as \( \lambda_p = 2\pi/q \gg r_0 \). Having \( \lambda_p \sim 700 \text{ nm} \), below we neglect this correction, and set \( n^2(\omega_p) = \epsilon_d + \langle \chi(\omega_p) \rangle \).

B. Global Phase Shift via Local Strong Coupling

In what follows we focus on the contribution of the ensemble of disordered single-molecule picocavities to the variation of the refractive index \( \Delta n(\omega_p) = \langle \chi(\omega_p) \rangle \), relative to a picocavity-free layer with dielectric constant \( \epsilon_d \). Detecting refractive index variations is standard in plasmonic sensing [60].

Due to the presence of picocavities, a probe wave that propagates in the metamaterial over a distance \( L \) is phase shifted by

\[ \Delta \Phi_L \equiv \text{Re}\{\Delta n(\omega_p)\} \omega_p L/c, \tag{8} \]

relative to propagation in the pure background dielectric. Since \( \text{Im}\{\epsilon_d\} \to 0 \) and \( |\langle \chi \rangle| \ll 1 \), we have that \( \Delta n(\omega_p) \approx \text{Re}\{\chi(\omega_p)\}/2 \). The susceptibility function scales as \( \langle \chi \rangle \sim (N/V)|d_{eg}|^2/2\hbar \), with \( N/V \) being the number density of single-molecule picocavities and \( |d_{eg}|^2 \) being the dipole moment for the transition induced at the probe frequency \( (d_{eg} = 3.8 \text{ D in Ref. [4]}) \). Large phase variations \( \Delta \Phi_L/\Phi_L \) of a few percent relative to the cavity-free background can be achieved with number densities \( N/V \sim 10 \mu \text{m}^{-3} \), which is consistent with our dilute regime assumptions and experiments [4, 61].

III. INTRACAVITY LIGHT-MATTER COUPLING SCHEME

Optical phase variations induced by the intracavity molecules depend on the dynamics of the internal molecular coherences, which determine the frequency dependence of \( \langle \chi(\omega_p) \rangle \). We study this dynamics for molecules within the lowest electronic potentials \( S_0 \) and \( S_1 \), as illustrated in Fig. 2c. The ground \((\nu = 0)\) and first excited \((\nu = 1)\) vibrational levels in \( S_0 \) are coupled to the lowest vibrational level \((\nu = 0) \) in \( S_1 \), by the probe and cavity fields, respectively. The Huang-Rhys factor [43] in \( S_1 \) is large enough to give a sizable oscillator strength for the \( \nu = 0 \leftrightarrow \nu = 1 \) vibronic sideband. The vibrational frequency \( \omega_v \) is assumed to exceed \( \hbar \omega_c = k_B T/ \text{h} \) at room temperature, as is typical with vinyl stretching modes \((\omega_v \approx 0.18 \text{ eV}) [35] \). The picocavity frequency is set to \( \omega_c = \omega_{00} - \omega_v \), with \( \omega_{00} \) being the \( 0 \rightarrow 0 \) vibronic absorption frequency. The cavity detuning from the \( \omega_{00} \) resonance ensures that the cavity field preferentially drives the \( 1 \rightarrow 0 \) transition, and in particular prevents driving population out of the ground vibrational state \((\nu = 0) \) in the absence of the probe.

In addition to the probe field at frequency \( \omega_p \) and the picocavity field at \( \omega_c \), we introduce in Fig. 2c an additional classical \emph{signal field} at frequency \( \omega_s \). The signal field drives the hot vibronic absorption band \( \nu = 1 \leftrightarrow \nu = 0 \). We show later that this signal field can be used as an optical switch, for controlling the molecular susceptibility at the probe frequency. In summary, we have the frequency hierarchy \( \omega_c < \omega_p < \omega_s \).

In order to compute \( \langle \chi(\omega_p) \rangle \), we label the relevant molecular transitions as \([1] \equiv |\nu = 0 \rangle, [2] \equiv |\nu = 1 \rangle, [3] \equiv |\bar{\nu} = 0 \rangle \), and \([4] \equiv |\bar{\nu} = 2 \rangle \), according to the scheme in Fig. 2c, to write a picocavity Hamiltonian of the form (we use \( \hbar = 1 \) throughout)

\[ \hat{H} = \omega_c \hat{a}^\dagger \hat{a} + \omega_{31}[2|2\rangle + \omega_{41}[3|3\rangle + \omega_{41}[4] \langle 4 \rangle + g_c[3|2\rangle \hat{a} + \Omega_p[3]\langle 1\rangle e^{-i\omega_s t} + \Omega_s[4]\langle 2\rangle e^{-i\omega_s t} + \mathcal{H.c.} \tag{9} \]
where \( \omega_{ij} = (E_i - E_j)/\hbar \) denote the molecular transition frequencies, \( g_c \) is the picocavity vacuum Rabi frequency, \( \Omega_p \) is the classical Rabi frequency of the probe field, and \( \Omega_s \) is the classical signal Rabi frequency. The bosonic cavity field operator is \( \hat{a} \), and \( \hat{H}_{\text{cav}} \) stays for Hermitian conjugation. Energy is given relative to the ground vibrational level (i.e., \( E_1 = 0 \)). Although the analysis reduces to an effective four-level system, the displaced oscillator picture for the \( S_0 \) and \( S_1 \) manifolds is helpful for anticipating potential issues in an experimental implementation of the scheme. It also serves as the basis of further studies that take energy transport dynamics into account [62].

For a quantized picocavity in the low-excitation manifold, the molecular basis should be supplemented with Fock states \( |n_c\rangle \) to give the dressed basis: \( |1\rangle \equiv |1; 0_c\rangle \), \( |2\rangle \equiv |2; 1_c\rangle \), \( |3\rangle \equiv |3; 0_c\rangle \), and \( |4\rangle \equiv |4; 1_c\rangle \). In the dressed-state picture, the probe field \( \Omega_p \) drives the transition \( |1\rangle \leftrightarrow |3\rangle \) within the vacuum manifold, and the signal field \( \Omega_s \) drives the transition \( |2\rangle \leftrightarrow |4\rangle \) within the one-photon manifold. The picocavity field admixes the vacuum and one-photon states \( |2\rangle \) and \( |3\rangle \). The coupling of the quantized picocavity field with other molecular transitions is neglected.

We model the evolution of the reduced density matrix \( \hat{\rho}(t) \) for an individual picocavity with a Lindblad quantum master equation of the form [42]

\[
\frac{d}{dt} \hat{\rho} = -i[\hat{H}, \hat{\rho}] + \sum_\alpha L_\alpha \hat{\rho} \hat{L}_\alpha^\dagger - \frac{1}{2}(L_\alpha^\dagger \hat{L}_\alpha \hat{\rho} + \hat{\rho} L_\alpha \hat{L}_\alpha^\dagger),
\]

where \( \hat{H} \) is given in Eq. (9), and \( \hat{L}_\alpha \) is the Lindblad operator associated with the \( \alpha \)-th dissipative channel. In Table I, we list the Lindblad operators used in this work, and the associated dephasing times in the bare basis. We use a notation in which the rate \( \gamma_{ij} \) describes the decay of the off-diagonal element \( \rho_{ij} \equiv \langle i|\hat{\rho}|j \rangle \), in the dressed basis. For example, vibrational relaxation from \( \nu = 1 \) (state \( |2\rangle \)) to \( \nu = 0 \) (state \( |1\rangle \)) in \( S_0 \) occurs at the rate \( \gamma_\nu \sim 1 \text{ ps}^{-1} \). On the other hand, the bare picocavity photon lifetime is \( \kappa^{-1} \sim 10 - 100 \text{ fs} \) [35]. Therefore, the combined decay rate of the dressed Raman coherence \( \langle 1|\hat{\rho}|2 \rangle \) is \( \gamma_{21} = \gamma_\nu/2 + \kappa/2 \), as \( |2\rangle \) has a single-photon character. The lifetime of the excited electronic state \( S_1 \) is \( \gamma_{31} \equiv \gamma_c \), given by either fluorescence or internal conversion. We set \( \gamma_{41} = \gamma_{31} \) throughout.

### IV. Homogeneous Autler-Townes Response

Starting from the master equation in Eq. (10), we follow Ref. [42] and derive a general expression for susceptibility for a homogeneous ensemble of \( N \) independent molecules in identical single-molecule picocavities, subject to classical driving by the probe and signal fields. We reproduce here the final expression, given by

\[
\chi(\omega_p) = \left( \frac{N}{V} \right) \frac{|d_{41}|^2}{\epsilon_0 \hbar} \times \frac{[\Delta_{21} + i \gamma_{21}] [\Delta_{41} + i \gamma_{41}] - \Omega_s^2}{[\Delta_{31} + i \gamma_{31}] [\Delta_{21} + i \gamma_{21}] [\Delta_{41} + i \gamma_{41}] - \Omega_s^2 - g_e^2 [\Delta_{41} + i \gamma_{41}]},
\]

and refer readers to Appendix B for the technical details of the derivation. In Eq. (11) we denote \( |d_{41}|^2 \) as the 0-0 oscillator strength, \( \Delta_{31} \equiv \omega_p - \omega_{31} \) is the probe detuning, \( \Delta_{21} \equiv \omega_p - \omega_c - \omega_{21} = \Delta_{31} - \Delta_c \) is the two-photon Raman detuning. Introducing the cavity detuning \( \Delta_c \equiv \omega_c - \omega_{32} \) and the signal detuning \( \Delta_s \equiv \omega_s - \omega_{42} \), we can write \( \Delta_{41} \equiv \Delta_{31} - \Delta_c + \Delta_s \). The ensemble susceptibility in Eq. (11) involves only the single-molecule coupling strength \( g_c \), highlighting the local character of the cavity-induced nonlinearity.

Although we focus on the polarization component at the probe frequency to derive Eq. (11), steady-state solutions for other polarization components at the cavity and signal field frequencies can also be derived from the Lindblad quantum master equation, resulting in a more general coupled mode theory that would describe the coherent interaction between different field components mediated by the material degrees of freedom [63, 64].
ing this more general problem is beyond the current scope of this work.

In Fig. 3, we plot the absorptive and dispersive parts of the disorder-free susceptibility \( \chi(\omega_p) \) around the bare 0-0 absorption resonance, under conditions of strong intracavity coupling. In Fig. 3a, we show the system response without the signal field (\( \Omega_s = 0 \)). The absorptive response shows two Autler-Townes (AT) peaks at \( \omega_p \approx \omega_{31} \pm g_c \). The doublet opens a broad semi-transparent window (solid line), due to cavity-induced AT splitting of the dressed states \( |3\rangle \) and \( |2\rangle \). The width of the Autler-Townes transparency window \( \Gamma_{AT} \) can be defined by \( A(\Gamma_{AT}/2) = A(g_c)/2, \) with \( A(\omega) = \text{Im}(\chi(\omega)) \). For a homogeneous system, \( \Gamma_{AT} \) scales linearly with the cavity coupling \( g_c \) [42]. On the other hand, the amount of residual absorption at the bare probe resonance (\( \Delta_p = 0 \)) can be shown to scale with the ratio \( \gamma_{21}/\gamma_{31} \) [65].

Figure 3a shows that at the center of the AT doublet, the probe field experiences normal dispersion (dashed line), in contrast to the “slow-light” dispersion expected for an interference-based transparency window [65]. The plot also shows a relatively broad region within the AT transparency window (order \( \gamma_{31} \) in frequency) in which probe dispersion overcomes absorption, i.e., \( \text{Re}\chi(\omega_p) > \text{Im}\chi(\omega_p) \). Clearly this condition always holds away from absorptive resonances (e.g., \( |\Delta_p| > 6\gamma_{31} \) in Fig. 3a). However, the ability of introducing a dispersive response in a frequency range that is otherwise opaque is a key resource for optical switching (see below).

In Fig. 3b, we show the susceptibility for the same conditions as in Fig. 3a, but now in the presence of a strong signal field (\( \Omega_s = \gamma_{31} \)), detuned from the 2 \( \rightarrow \) 4 transition frequency (\( \Delta_s = \gamma_{42} \)). The positions of the peaks AT \( \pm \) remain practically unchanged, and the probe response continues to exhibit the same AT doublet. However, now a two-photon absorption resonance (TPA\(_{41}\)) destroys the transparency window: we see a broad background with a peak at the TPA\(_{41}\) resonance condition, \( \Delta_p - \Delta_s + \Delta_c = 0 \). For \( \Delta_s = 0 \) and \( \Delta_c = \gamma_{31} \), the resonance occurs at \( \Delta_p = -\gamma_{31} \). This TPA channel can be understood as a result of the mixing between the dressed states \( |2\rangle \) and \( |3\rangle \), mediated by an intracavity vacuum that acts as an effective doorway mechanism for the \( (\omega_p, \omega_p) \) process:

\[
|1\rangle|0_c\rangle \rightarrow \omega_p \leftrightarrow \langle 3|0_c\rangle \leftrightarrow \langle 2|1_c\rangle \rightarrow \omega_p \leftrightarrow \langle 4|1_c\rangle.
\]

Note that the classical probe and signal fields do not change the intracavity photon number. This emerging TPA channel effectively modulates the dispersive response of the probe field within the AT window, by making absorption comparable to the dispersion (Fig. 3b, shaded area). As we discuss below, at lower signal beam intensities the detrimental effect of the signal field can be small enough to preserve its dispersive properties in spite of increased absorption. Additional nonlinear absorption channels such as hot absorption \( 1 \rightarrow 1 \) can be suppressed, either by controlling the probe field intensity or selecting molecular vibrations with different fundamental frequencies in \( S_0 \) and \( S_1 \) (e.g., azobenzene [66]).

V. DISORDER-AVERAGED AUTLER-TOWNES RESPONSE

A. Rabi and energy disorder

In this section, we study the lineshape of the AT transparency window under two types of structural disorder: random intracavity volumes and random molecu-
lar transition frequencies. The first arises from the distribution of gap volumes $V_g$ in the picocavity ensemble. The volume distribution is assumed to be Gaussian, with mean value $\langle V_g \rangle$ and standard deviation $\sigma_{V_g}$. Given that $g_c = f/\sqrt{V_g}$ \cite{67}, with $f$ a constant, the distribution of Rabi frequencies $g_c$ is also Gaussian with mean value $\langle g_c \rangle = f/\sqrt{\langle V_g \rangle}$ and variance $\sigma_{g_c}^2 = f^2 \sigma_{V_g}^2 / 2 \langle V_g \rangle^{3/2}$, where only leading terms in the small parameter $\sigma_{V_g}/\langle V_g \rangle$ are kept.

In Fig. 4, we illustrate the impact of Gaussian energy disorder of the molecular transition frequencies. The inhomogeneous width of the cavity-free probe absorption band is $\sigma_{31}$ (FWHM $\approx 2.4 \sigma_{31}$). The ground state vibrational band $\nu = 0 \rightarrow \nu = 1$ has inhomogeneous width $\sigma_{21}$, and width of the hot-band absorption $\nu = 1 \rightarrow \nu = 2$ is $\sigma_{32}$. Since pure vibrational linewidths ($\sigma \sim 4$ meV \cite{68}) are much smaller than typical vibronic linewidths ($\sigma \sim 100$ meV \cite{43}), we have $\sigma_{21}/\sigma_{31} \ll 1$ and $\sigma_{32} \sim \sigma_{31}$.

In Fig. 5a, we plot the AT absorption doublet for an ensemble with a Gaussian distribution of Rabi frequencies, but otherwise homogeneous, in the absence of a signal field ($\Omega_s = 0$). In comparison with the fully homogeneous response from the previous section, the doublet lineshape remains largely unaltered even for broad distributions with $\sigma_{g_c} \approx \langle g_c \rangle$. This is reminiscent of the weak dependence of the cavity response on the distribution of dipole moment orientations found in Ref. \cite{42}. Therefore we neglect both orientational and mode volume disorder in what follows.

In Fig. 5b, the AT lineshape is shown for a fixed Rabi frequency ($\sigma_{g_c} \rightarrow 0$), but the molecular levels are inhomogeneously broadened, and the response is numerically averaged over a Gaussian distribution of energy levels. No signal field is applied. We see that the overall doublet shape of the AT transparency window is insensitive to the increase of the vibronic linewidth $\sigma_{31}$. This is expected, as the AT transparency window is sustained by the cavity-induced Raman coherence $\rho_{21} = (1, 0, |1\rangle, |2\rangle, 1_c)$. Therefore, the photon decay rate $\kappa$.

B. Theoretical limits for phase modulation

We now study the feasibility of the proposed optical switch in disordered picocavity ensembles. In order for the relative phase shift $\Delta \Phi_L/\Phi_L$ in Eq. (8) to be detectable, dispersion of the probe field should overcome...
absorptive losses. Therefore, we use the figure-of-merit
\[
\eta_p(\omega_p) \equiv \frac{\text{Re} \chi(\omega_p)}{\text{Im} \chi(\omega_p)},
\]
to quantify the theoretical performance of the metamaterial for phase modulation at the probe frequency. The global dispersive behavior of the medium (phase shifting) prevails over inherent molecular absorption losses when \(\eta_p\) exceeds unity over a bandwidth \(\gamma_{31}\). In a heterodyne setting that detects the interference of a transmitted probe field \(E_p(\omega_p)\) with a reference beam, the ratio \(\eta_p\) correlates with the fringe visibility. If the transmitted probe field is largely attenuated (\(\eta_p \ll 1\)), interference with a reference field cannot be resolved.

To account for energy disorder, we average Eq. (12) numerically using independent Gaussian distributions for each molecular transition frequency. We also estimate this average analytically using independent Lorentzian distributions for the molecular transition frequencies. As we prove in Appendix C, the Lorentzian averaging reduces to replacing \(\gamma_{ij}\) in Eq. (11) with \(\Sigma_{ij} = \gamma_{ij} + \sigma_{ij}\) everywhere. These analytical results allow us to gain insight into the multiple parameters that determine the effective nonlinear probe response, and we confirm numerically that a more realistic Gaussian disorder gives the same trends for key observables as the Lorentzian disorder model. The Lorentzian disorder technique has also been used in Refs. [42, 69, 70] to simplify the average of system observables over a random distribution of Hamiltonian parameters.

We can rewrite Eq. (12) for the figure-of-merit by separating the real and imaginary parts of the susceptibility (11). As a next step, we apply our technique of averaging over Lorentzian disorder, developed in Ref. [42] and described in detail in Appendix C. In the absence of a signal field (\(\Omega_s = 0\)), the ratio \(\eta_p\) as a function of probe detuning \(\Delta_p\) for a resonant cavity (\(\Delta_c = 0\)) becomes:
\[
\eta_p(\Delta_p) = \frac{g_c^2 \Delta_p - \Delta_p(\Delta_p^2 + \Sigma_{21}^2)}{\Sigma_{31}(\Delta_p^2 + \Sigma_{21}^2) + \Sigma_{21}^2 g_c^2},
\]
where the parameters \(\Sigma_{ij} \equiv \gamma_{ij} + \sigma_{ij}\) represent total decoherence rates, including homogeneous (\(\gamma_{ij}\)) and inhomogeneous (\(\sigma_{ij}\)) contributions. For the numerical averaging over Gaussian disorder, we used this expression with \(\Sigma_{ik} \rightarrow \gamma_{ik}\).

In a cavity-free scenario, Eq. (13) reduces to the linear scaling \(\eta_p = -\Delta_p/\Sigma_{31}\). This linear dependence is reproduced also if the averaging over inhomogeneous broadening is carried out numerically using Gaussian disorder (shown as a dotted line in Fig. 6a). The cavity vacuum induces a deviation from this linear scaling. As numerical averaging shows (solid and dashed lines in Fig. 6a), the figure-of-merit increases and exhibits a maximum when the probe is slightly detuned from the center of the AT window, with the maximum value \(\eta_{\text{max}}\) considerably exceeding 1. We can estimate the optimal detuning at which \(\eta_{\text{max}}\) is reached using our analytical Lorentz-averaged model (13). We find:
\[
\Delta_{\text{p, optimal}} \approx g_c \sqrt{\frac{\Sigma_{21}}{\Sigma_{31} + 3\Sigma_{21}}},
\]
where we dropped quadratic terms in \(\Sigma_{21}/g_c\). For low-quality picocavities in strong coupling regime, we have \(\gamma_c \ll \kappa \ll \gamma_{31} < g_c\).

We now can estimate \(\eta_{\text{max}} = \eta_p(\Delta_{\text{p, optimal}})\). In the framework of the Lorentzian disorder model, the maximum figure-of-merit \(\eta_{\text{max}}\) is:
\[
\eta_{\text{max}} = \frac{g_c}{\sqrt{\Sigma_{21}(\Sigma_{31} + 3\Sigma_{21})}} \left( \frac{\Sigma_{31} + 2\Sigma_{21}}{2\Sigma_{31} + 3\Sigma_{21}} \right),
\]
where we have ignored terms that are second order in $\Sigma_{21}/g_c$. If $\Sigma_{21}$ is small, then $\eta_{\text{max}}$ is large, which is the case we studied in Ref. [42]. Consider now an ensemble of lossy picocavities with a large Raman decoherence rate $\Sigma_{21} \sim \Sigma_{31}$. Equation (15) then predicts that cavity-mediated optical phase modulation within the AT transmission window can still be feasible, provided that the single-molecule Rabi coupling is strong enough. For example, we can achieve $\eta_{\text{max}} \geq 1$ even for $\Sigma_{21}/\Sigma_{31} = 1$, with Rabi couplings $g_c/\Sigma_{31} \geq 3.33$. For a representative zero-phonon width $\Sigma_{31} \approx 50$ meV [4, 12], this corresponds to $g_c \approx 167$ meV. Improving the quality factor of the picocavities ($\Sigma_{31} \sim \kappa$), such that the ratio $\Sigma_{21}/\Sigma_{31}$ decreases by a factor of two, reduces the constraint on single-molecule coupling to $g_c \geq 98$ meV, for phase modulation to be detectable. Single-molecule couplings of these magnitudes are within experimental reach [1, 3, 4, 71].

Let us now check if the Lorentz approximation is consistent with numerical Gaussian-based approach. For the system parameters used in Fig. 6a, the Lorentz disorder model predicts the maximum figure-of-merit $\eta_{\text{max}} \approx 2.0$ at $\Delta_p = 0.55 \gamma_{31}$. This detuning is only slightly higher than the value $\eta_{\text{max}} = 1$, predicted by numerically averaging Eq. (13), with $\Sigma_{ik} \rightarrow \gamma_{ik}$, over independent Gaussian frequency distributions (Fig. 6a, solid line). In the presence of a signal field ($\Omega_s > 0$), blue-detuned from the 4-2 resonance by $\Delta_s = \gamma_{13}$, the AT window lineshape becomes distorted (see Fig. 3), and $\eta_{\text{max}}$ decreases monotonically with increasing $\Omega_s$ (Fig. 6a, dashed line), as discussed in more detail in Section V C.

Complementary to the discussion of the frequency-dependent phase performance parameter $\eta_p(\omega_p)$, we can use the Lorentz disorder model to analyze the lineshape of the Autler-Townes window. As already mentioned, the position of the AT$^+$ doublet peaks is largely insensitive to disorder. However, transparency within the AT window is reduced as the quality of the vibrational Raman coherence $\rho_{21}$ degrades with increasing $\gamma_{21}$ and $\sigma_{21}$, which decreases the width $\Gamma_{\text{AT}}$.

In the Lorentz disorder model, the AT width can be written as

$$\Gamma_{\text{AT}} \approx 2\sqrt{g_c^2 - \sqrt{2}g_c(\Sigma_{31} + \Sigma_{21})}, \quad (16)$$

which shows that the AT window is formally closed when $g_c \leq \sqrt{2}(\Sigma_{21} + \Sigma_{31})$.

Figure 6b shows that the Lorentz disorder model in general overestimates $\Gamma_{\text{AT}}$ relative to Gaussian averaging. For the parameters in Fig. 6b, the AT window is predicted by Eq. (16) to close for $\gamma_{21} = 1.78 \gamma_{31}$, which agrees well with the value ($1.71 \gamma_{31}$) obtained for Gaussian disorder. Figure 6b also shows that Eq. (15) correctly captures the scaling of $\eta_{\text{max}}$ with the Raman decay rate $\gamma_{21}$, for narrow vibrational coherences with $\sigma_{21} \ll \sigma_{31}$.

We conclude that $\eta_p > 1$, which correspond to observable phase shifts, can be achieved in our metamaterial in spite of strong energy disorder and ultra-fast cavity photon decoherence, as long as the coupling constant $g_c$ is large enough.

C. Controlling the probe phase shift with a weak signal laser

Let us finally discuss the nonlinear interaction between the probe field at $\omega_p$ and an additional signal laser at $\omega_s$, mediated by the ensemble of single-molecule picocavities. In the presence of the signal field, we derive from Eq. (11) a more general expression for $\eta_p$ than Eq. (13), which captures the dependence of $\chi_2$ on the signal frequency $\omega_s$ and its Rabi frequency $\Omega_s \propto |\mathbf{E}_s|$. We then average $\eta_p$ over independent fluctuations of the molecular transition frequencies $\omega_{31}, \omega_{32}$ and $\omega_{42}$. Assuming the Lorentzian disorder model and applying the Lorentz averaging technique outlined in the Appendix C, we obtain a disorder-averaged expression for $\eta_p$ that reads [42]

$$\eta_p(\Delta_p) = g_c^2(\Delta_p - \lambda_p \Delta_{41}) - \Delta_p[(\Delta_p - \lambda_p \Delta_{41})^2 + (\Sigma_{21} + \lambda_p \Sigma_{41})^2] + \frac{\Sigma_{31}(\Delta_p - \lambda_p \Delta_{41})^2 + (\Sigma_{21} + \lambda_p \Sigma_{41})^2 + g_c^2(\Sigma_{21} + \lambda_p \Sigma_{41})}{(\Sigma_{31})^2}, \quad (17)$$

where $\Delta_{31} = \Delta_{21} = \Delta_p$. This expression can be used for numerical modeling with Gaussian disorder upon standard replacement $\Sigma_{ik} \rightarrow \gamma_{ik}$. Here we have introduced...
the dimensionless signal parameter
\[ \lambda_s = \frac{\Omega_s^2}{(\Delta_{41}^2 + \Sigma_{41}^2)}. \]  

such that Eq. (17) reduces to Eq. (13) when \( \lambda_s = 0 \). The impact of the signal field is determined entirely by \( \lambda_s \). We find that finite \( \lambda_s \) result in suppression of \( \eta_p \), so that the \textit{probe-only} performance in Eq. (13) corresponds to the upper bound of performance of the metamaterial with both probe and signal fields being present. Such a suppression is explained by the observation that the signal field tends to close the AT transparency window (see Figure 3b). Fortunately, reaching \( \eta_p > 1 \) is still quite possible with small values of \( \lambda_s \). For example, the dashed line in Fig. 6a corresponds to the signal-on arrangement with \( \lambda_s = 0.036 \). As Eq. (18) shows, small values of \( \lambda_s \) are achieved either by reducing the strength of the signal field, or by detuning it from the \( |2 \rangle \rightarrow |4 \rangle \) transition (since \( \Delta_{41} = \Delta_p + \Delta_s \)). Both these actions effectively reduce the detrimental effect of losses via the two-photon absorption channel TPA_{41}.

Our metamaterial thus implements a weak-field controllable optical nonlinearity, with the phase shift of the probe beam being controlled by the strength of the signal beam. It can also be used as an optical switch; when \( \eta_p > 1 \), the probe beam passing through the disordered metamaterial preserves coherence and can interfere with the reference beam, whereas \( \eta_p < 1 \) means loss of coherence and destroyed interference. The crossover between these two regimes is reversible and is controlled by simply changing the intensity of the signal beam.

In Fig. 7, we show a parameter map \((g_c, \Omega_s, \eta_{\text{max}})\) for optical switching with a signal field detuned by \( \Delta_s = 1.0 \gamma_{31} \). We assume a relatively large Raman decay rate \( \gamma_{21} = 0.5 \gamma_{31} \), to highlight the feasibility of the optical switching scheme under realistic picocavity conditions. For a typical \( S_1 \) radiative lifetime \( 1/\gamma_{31} \sim 1 \) ns and inhomogeneous width \( \sigma_{31} \sim 50 \) meV, Fig. 7 shows that signal pulses with far-field intensities \( I_s \sim 10 \text{ kW/cm}^2 \) (\( d_{32} \sim 1 \) D), are sufficient to switch off coherent phase modulation, in an ensemble of resonant picocavities in the Autler-Townes regime \((g_c > \Sigma_{31})\). These field intensities are orders of magnitude smaller than the typical two-photon excitation intensities used in photochemistry \((\sim 10^6 \text{ W/cm}^2 \) [72, 73]) or microscopy \((I \sim 10^{15} \text{ W/cm}^2 \) [74, 75]), which can facilitate the implementation of an optical switch based on the Autler-Townes window.

VI. CONCLUSIONS

We perform a proof-of-principle theoretical analysis of coherent optical phase manipulation assisted by the electromagnetic vacuum in a dilute ensemble of disordered single-molecule plasmonic picocavities. We show that strong light-matter coupling with individual organic chromophores that have well-resolved vibronic progressions in the absorption and emission spectra opens an Autler-Townes transparency band [65] in the \( 0 \rightarrow 0 \) sideband of the chromophore absorption spectrum, which we propose to use for coherent manipulations of the average refractive index of the disordered medium at these frequencies, resulting in a controllable phase shift of a propagating probe wave tuned on resonance with the \( 0 \rightarrow 0 \) sideband. Although the achievable refractive index variations may be small in a potential realization of the proposed scheme, plasmonic nanocavity structures can in principle be engineered to detect refractive index changes at optical frequencies of 0.1% or less [76].

Assuming a Lorentzian model for the picocavity spectrum and a displaced-oscillator model for the chromophore levels [17, 18, 29], we obtain an analytical expression for the ratio between the dispersive and absorptive parts of the average system susceptibility at a probe frequency, which highlights the dependence of the predicted phase shift on important design parameters such as the inhomogeneous linewidth of the relevant vibronic transitions, the radiative and non-radiative molecular relaxation rates, the picocavity photon lifetime, and the average single-molecule Rabi frequency. The predicted phase signal may be challenging to detect with current plasmonic picocavities [1–4], but expected improvements in nanofabrication may enable the observation of the proposed phase control scheme.

In our analysis we take into account realistic sources of disorder and relaxation including a distribution of Rabi couplings, energy disorder in the molecular transition frequencies, vibrational relaxation and photon losses. The figure-of-merit for the proposed picocavity-induced phase shift at the probe frequency was found to be significantly more sensitive to static disorder in the molecular transition frequencies than disorder in the Rabi coupling strength.

The proposed phase shift of the probe field can be dynamically gated with an additional signal field at a higher frequency set to drive an excited state molecular coherence. The phase switch mechanism is interpreted as the opening of a novel two-color two-photon absorption process in the molecules, mediated and enhanced by the ensemble of picocavity vacua. Signal fields as weak as 10 kW/cm² are estimated to be sufficient for implementing optical switching behavior in a disordered picocavity ensemble. Our work thus opens the way toward the development of few-photon nonlinear optical devices with molecular picocavity metamaterials.

VII. ACKNOWLEDGMENTS

F.H. is funded by ANID – Fondecyt Regular 1181743 and Millennium Science Initiative Program ICN17_012.
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### Appendix A: Derivation of the Effective Index at the Probe Frequency

Here we derive Eq. (6) from the main text, following the procedure created in Refs. [54, 55] for various kinds of random media that allow for perturbative approach. Our starting point is Eq.(1). Its $n$-th discrete frequency component oscillating at the frequency $\omega_n$ satisfies the wave equation:

$$\nabla^2 E_n(r) + \frac{\epsilon_0 \omega_n^2}{\epsilon^2} E_n(r) = -\frac{\omega_n^2}{\epsilon_0 \epsilon^2} P_n(r), \quad (A1)$$

where we used $D_n(r) = \epsilon_0 \epsilon_d E(r)$, and $P_n(r)$ is the component of the polarization density at $\omega_n$, caused by inclusions (cavities with embedded point dipoles). We assume that at the probe frequency ($\omega_p = \omega_p$) a linear relationship between the polarization and the (weak) probe field holds, so that

$$P_p(r) = \epsilon_0 \chi L(\omega_p, r) \cdot E_p(r), \quad (A2)$$

where the susceptibility $\chi = \chi(\omega, r)$ captures the fact that the polarization is created locally, and vanishes between the picocavities.

Following the effective medium approach from Refs. [54–58], we write the electric field and the susceptibility as

$$E_p(r) = \langle E_p(r) \rangle + \delta E_p(r), \quad (A3)$$

$$\chi(r) = \langle \chi \rangle + \delta \chi(r), \quad (A4)$$

where $\langle E_p(r) \rangle$ is the average probe field that propagates according to the effective index $n(\omega_p)$, and $\delta E_p(r)$ is a position-dependent fluctuation of the electric field caused by the presence of picocavities; $\langle \chi \rangle$ is the uniform effective susceptibility of the medium (the effective medium correction), and $\delta \chi(r)$ is the local fluctuation of the response. Our mean values are chosen in such a way that, by construction,

$$\langle \delta E_p(r) \rangle = \langle \delta \chi(r) \rangle = 0. \quad (A5)$$
Inserting Eqs. (A2), (A3), and (A4) into Eq. (A1), and averaging the result, we obtain an equation for the averages, which reads:

\[ \nabla^2 \langle \mathbf{E}(r) \rangle + \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] \langle \mathbf{E}(r) \rangle = -\frac{\omega_p^2}{\epsilon_0 c^2} \langle \delta \chi(r) \delta \mathbf{E}(r) \rangle, \]

(A6)

where we used (A5) to eliminate the terms proportional to \( \delta \chi \) and \( \delta \mathbf{E} \) times a position-independent factor. We then subtract (A6) from our starting-point equation [Eq.(A1) with (A2), (A3), and (A4) plugged in]. This leaves us with an equation for the fluctuations:

\[ \nabla^2 \delta \mathbf{E}(r) + \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] \delta \mathbf{E}(r) = -\frac{\omega_p^2}{\epsilon_0 c^2} \delta \chi(r) \langle \mathbf{E}(r) \rangle - \frac{\omega_p^2}{\epsilon_0 c^2} [\delta \chi(r) \delta \mathbf{E}(r) - \langle \delta \chi(r) \delta \mathbf{E}(r) \rangle]. \]

(A7)

For now we will assume that the role of the fluctuations is relatively small; we will quantify this assumption below. This allows us to neglect the second-order term \( \propto \delta \chi(r) \delta \mathbf{E}(r) - \langle \delta \chi(r) \delta \mathbf{E}(r) \rangle \) in the last equation. Our position-dependent first-order equations describing fields in the disordered medium thus become:

\[ \nabla^2 \langle \mathbf{E}(r) \rangle + \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] \langle \mathbf{E}(r) \rangle = -\frac{\omega_p^2}{\epsilon_0 c^2} \delta \chi(r) \delta \mathbf{E}(r), \]

(A8)

\[ \nabla^2 \delta \mathbf{E}(r) + \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] \delta \mathbf{E}(r) = -\frac{\omega_p^2}{\epsilon_0 c^2} \delta \chi(r) \langle \mathbf{E}(r) \rangle. \]

(A9)

We keep the second-order term in the equation (A8) for the averages, since there it is the lowest-order disorder-dependent contribution.

The next step is to Fourier-transform these equations. This gives:

\[ \left( \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] - k^2 \right) \langle \mathbf{E}(k) \rangle = -\frac{\omega_p^2}{\epsilon_0 c^2} \int dq \langle \delta \chi(k - q) \delta \mathbf{E}(q) \rangle; \]

(A10)

\[ \left( \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] - k^2 \right) \delta \mathbf{E}(k) = -\frac{\omega_p^2}{\epsilon_0 c^2} \int dq' \langle \delta \chi(k - q') \delta \mathbf{E}(q') \rangle. \]

(A11)

We solve Eq.(A11) for \( \delta \mathbf{E}(k) \), and plug it into the integrand in Eq.(A10). We get:

\[ \left( \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] - k^2 \right) \langle \mathbf{E}(k) \rangle = \left( \frac{\omega_p^2}{\epsilon_0 c^2} \right)^2 \int dq \int dq' \langle \mathbf{E}(q') \rangle \frac{\langle \mathbf{E}(q') \rangle}{\left( \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] - q^2 \right)^2} \langle \delta \chi(k - q) \delta \chi(q - q') \rangle. \]

(A12)

This integral equation for \( \langle \mathbf{E}(k) \rangle \) shows that the field average depends on two deterministic components: the propagator of electromagnetic waves in the effective medium, \( G(q) = \left( \frac{\omega_p^2}{c^2} [\epsilon_d + \langle \chi \rangle] - q^2 \right)^{-1} \), and the correlator \( \langle \delta \chi(k - q) \delta \chi(q - q') \rangle \). We can simplify it further by examining the properties of this correlator. Let us denote as

\[ K(|r_1 - r_2|) = \langle \delta \chi(r_1) \delta \chi(r_2) \rangle \]

(A13)

the spatial correlator of the susceptibility fluctuations. This form is very general and only assumes that the medium is homogeneous and isotropic on average (and hence the correlator depends only on \( |r_1 - r_2| \)). We can now write:

\[ \langle \delta \chi(k - q) \delta \chi(q - q') \rangle = \frac{1}{(2\pi)^6} \int dr_1 dr_2 e^{-i(k-q)\cdot r_1} e^{-i(q-q')\cdot r_2} K(|r_1 - r_2|) \]

(A14)

\[ = \frac{1}{(2\pi)^6} \int dr_2 e^{-i(k-q')\cdot r_2} \int d(|r_1 - r_2|) e^{-i(k-q)(r_1-r_2)} K(|r_1 - r_2|) \]

(A15)

\[ = \delta(k - q') K(|k - q|) \]

(A16)

with

\[ K(q) = \frac{1}{(2\pi)^3} \int dr \ e^{-iqr} K(r) \]

(A17)
being the Fourier transform of the spatial correlator.

We now plug Eq. (A16) into Eq. (A12) and carry out integration over \( q' \). Thanks to the delta-function, \( \langle E(q') \rangle \) transforms into \( \langle E(k) \rangle \) and cancels out in the both sides of this equation. This happens since the system is homogeneous on average. Therefore, instead of an integral equation involving Fourier components of the field amplitude, we are left simply with the following modified dispersion equation of electromagnetic waves in our disordered medium:

\[
\left( \frac{\omega_p^2}{c^2} \kappa + (\chi) \right) - K^2 = \left( \frac{\omega_p^2}{\epsilon_0 c^2} \right)^2 \int dq \, G(q) K(|k - q|) \equiv \frac{\omega_p^2}{c^2} \Delta \epsilon_p, \tag{A18}
\]

where \( \Delta \epsilon_p = \omega_p^2/(c^2 \epsilon_0^2) \). \( \int dq \, G(q) K(|k - q|) \) is a deterministic, coordinate-independent correction to the averaged susceptibility \( \langle \chi \rangle \).

The discussion in the text is done in the effective medium approximation, under the assumption that \( \Delta \epsilon_p \) is negligibly small. Eq. (A18) allows one to verify this assumption for a given set of parameters relevant to their system, and to quantify the error associated with this assumption. The full procedure requires (i) introducing a sample-specific Green’s function from the integration axis, and (iii) carrying the integration out explicitly. Below we will only show that having this correction small is equivalent to the requirement of having a wavelength, \( \lambda \), large in comparison with the average scale \( r_0 \), at which the fluctuations happen.

Consider, for example, Gaussian spatial correlations: \( K(r) = K_0 \, e^{-r^2/(2r_0^2)} \), where the constant \( K_0 \) expresses the magnitude of the correlator, and \( r_0 \) is the scale of the order of the inclusion size; by choosing it this way we basically say that the correlations between the electric field and induced susceptibility vanish outside of each picocavity. Then the magnitude of its Fourier transform \( K(q) = K_0 \, e^{-q^2 r_0^2/2} \) is indeed determined by the ratio \( r_0/\lambda^2 \) in the exponent, which vanishes as long as for the typical wavelength \( \lambda = 2\pi/q \gg r_0 \). Based on the analysis of Ref. [58], that shows that the results of this approach are not sensitive to the specific choice of the correlator, we can argue that it is a general, correlator-independent property of our metamaterial.

Appendix B: Derivation of the Effective Probe Susceptibility

Here we outline the steps in the derivation of Eq. (11) in the main text, following closely the method used in Ref. [42]. In order to describe light-matter interaction in an effective four-level vibronic state manifold, the molecular system Hamiltonian \( \hat{\mathcal{H}} \) from Eq. (9) is used in a Lindblad quantum master equation [Eq. (10)], which reads

\[
\frac{d}{dt} \hat{\rho} = -i [\hat{\mathcal{H}}, \hat{\rho}] + \mathcal{L}_\kappa [\hat{\rho}] + \mathcal{L}_{\gamma_c} [\hat{\rho}] + \mathcal{L}_{\gamma_{\perp}} [\hat{\rho}] + \mathcal{L}_{\gamma_r} [\hat{\rho}] \tag{B1}
\]

with the Lindblad operators \( \mathcal{L} \) having the decay timescales listed in Table I. \( \mathcal{L}_\kappa [\hat{\rho}] \) describes photon decay within the picocavity, with \( \kappa \sim 10 - 100 \, \text{fs}^{-1} \) being the fastest decay timescale in the problem. \( \mathcal{L}_{\gamma_c} [\hat{\rho}] \) and \( \mathcal{L}_{\gamma_{\perp}} [\hat{\rho}] \) correspond to intramolecular vibration-assisted relaxation within the potentials \( S_0 \) to \( S_1 \), respectively, for decay times in the picosecond regime. The term \( \mathcal{L}_{\gamma_r} [\hat{\rho}] \) describes the decay of the lowest electronic singlet excitation at rate \( \gamma_r \). The corresponding dissipators are given by:

\[
\mathcal{L}_\kappa [\hat{\rho}] = (\kappa/2) \{ 2\hat{a} \hat{\rho} \hat{a}^\dagger - \hat{a}^\dagger \hat{a} \hat{\rho} - \hat{\rho} \hat{a} \hat{a}^\dagger \}, \\
\mathcal{L}_{\gamma_c} [\hat{\rho}] = (\gamma_{c}/2) \{ 2|1\rangle \langle 2| \hat{\rho} |2\rangle \langle 1| - |2\rangle \langle 2| \hat{\rho} - \hat{\rho} |2\rangle \langle 2| \}, \\
\mathcal{L}_{\gamma_{\perp}} [\hat{\rho}] = (\gamma_{\perp}/2) \{ 2|3\rangle \langle 4| \hat{\rho} |3\rangle \langle 4| - |4\rangle \langle 4| \hat{\rho} - \hat{\rho} |4\rangle \langle 4| \}, \\
\mathcal{L}_{\gamma_r} [\hat{\rho}] = (\gamma_{r}/2) \{ 2|1\rangle \langle 3| \hat{\rho} |3\rangle \langle 1| - |3\rangle \langle 3| \hat{\rho} - \hat{\rho} |3\rangle \langle 3| + 2|2\rangle \langle 3| \hat{\rho} |3\rangle \langle 2| - |3\rangle \langle 3| \hat{\rho} - \hat{\rho} |3\rangle \langle 3| \}. \tag{B2}
\]

In order to obtain Eq. (11), we focus on matrix elements \( \hat{\rho} \) that explicitly account for the presence or absence of the cavity photon and introduce the notation \( \rho_{ij}^{n}(t) = \langle i; m_c| \hat{\rho}(t)| j; n_c \rangle \), where \( |i \rangle \) and \( |j \rangle \) represent molecular states \( i, j = 1, 2, 3, 4 \), and \( |m_c \rangle \) and \( |n_c \rangle \) represent cavity Fock states with photon numbers \( m_c \) and \( n_c \), respectively. In order to remove fast oscillations from the equations of motion, we define slowly-varying amplitudes \( \sigma_{ij}^{n} \) for selected elements of reduced density matrix as follows: \( \sigma_{00}^{00} = \rho_{00}^{00} e^{-i \omega_p t} \), \( \sigma_{01}^{01} = e^{-i \omega_p t} \rho_{12}^{01} \), \( \sigma_{02}^{01} = \rho_{22}^{01} \), \( \sigma_{04}^{01} = e^{-i \omega_p t} \rho_{43}^{01} \), and \( \sigma_{11}^{01} = e^{-i \omega_p t} \rho_{11}^{01} \). In terms of these slowly-varying amplitudes, we obtain from Eq. (B1) the following equations of motion for the coherences:
\[ \dot{\sigma}_{13}^{00} = i(\omega_{41} - \omega_p)\sigma_{13}^{00} - \gamma_{31}\sigma_{13}^{00} - i\Omega_p(\sigma_{33}^{00} - \sigma_{11}^{00}) + ig_c\sigma_{12}^{01} \]
\[ \dot{\sigma}_{12}^{01} = i(\omega_{21} + \omega_c - \omega_p)\sigma_{12}^{01} - \gamma_{21}\sigma_{12}^{01} - i\Omega_p\sigma_{32}^{01} + ig_c\sigma_{13}^{00} + i\Omega_s\sigma_{14}^{01} \]
\[ \dot{\sigma}_{32}^{01} = -i(\omega_{32} - \omega_c)\sigma_{32}^{01} - \gamma_{32}\sigma_{32}^{01} - ig_c(\sigma_{12}^{11} - \sigma_{33}^{00}) - i\Omega_p\sigma_{12}^{01} + i\Omega_s\sigma_{34}^{01} \]
\[ \dot{\sigma}_{14}^{01} = i(\omega_{41} + \omega_c - \omega_p - \omega_s)\sigma_{14}^{01} - \gamma_{41}\sigma_{14}^{01} - i\Omega_p\sigma_{34}^{01} + i\Omega_s\sigma_{12}^{01} \]
\[ \dot{\sigma}_{34}^{01} = i(\omega_{43} + \omega_c - \omega_s)\sigma_{34}^{01} - \gamma_{43}\sigma_{34}^{01} - i\Omega_p\sigma_{14}^{01} - ig_c\sigma_{22}^{11} + i\Omega_s\sigma_{32}^{01} \]
\[ \dot{\sigma}_{24}^{11} = i(\omega_{42} - \omega_s)\sigma_{24}^{11} - \gamma_{42}\sigma_{24}^{11} - ig_c\sigma_{34}^{01} + i\Omega_s(\sigma_{22}^{11} - \sigma_{44}^{11}) \]

where we introduced the decay rates \( \gamma_{31} = \gamma_c/2, \gamma_{21} = \kappa/2 + \gamma_c/2, \gamma_{32} = \kappa/2 + \gamma_c/2, \gamma_{43} = \kappa/2 + \gamma_c/2, \gamma_{42} = \kappa + \gamma_c/2, \) and \( \gamma_{41} = \gamma_{43} \). The homogeneous probe linewidth is \( \gamma_{31} \) and the Raman linewidth is \( \gamma_{43} \). In deriving Eqs. (B3)-(B3), we neglect the contribution of states such as \( |2, 0_c, \rangle \), \( |3, 1_c, \rangle \), or \( |4, 0_c, \rangle \), which are neither populated nor driven under our imposed assumptions of stationarity and weak signal and probe driving. Accounting for such states would result, for example, in the addition of an extra term proportional to \( \sigma_{13}^{01} \) in the right-hand side of equation (B3) for \( \dot{\sigma}_{13}^{00} \), term that can be shown to vanish in the stationary limit. In other words, the set of Eqs. (B3) does not correspond to a complete description of the system coherences, but can be considered as a minimal set of equations of motion that can account for the non-linear optical response of our system of interest. The homogeneous probe susceptibility \( \chi_p \) is then obtained by algebraically solving for the steady state probe coherence \( \sigma_{13}^{00}(t \to \infty) \) from the coupled system of equations (B3), using the relation \( \chi_p = \sigma_{13}^{00}(\infty)/E_p \), which gives Eq. (11).

Appendix C: Averaging over Energy Disorder: The Lorentzian Technique

Assume that the energies of the states \( |1, \rangle, |2, \rangle, |3, \rangle \) and \( |4, \rangle \) fluctuate as a result of structural disorder (random environment). Then the detunings \( \Delta_{31} = \omega_p - \omega_3 + \omega_1 \) and \( \Delta_{21} = \omega_p - \omega_2 + \omega_1 \) also become random quantities. To find the disorder-averaged response, we can numerically integrate Eq. (11) in the main text. However, here we will explore an alternative route. Instead of using a realistic Gaussian distribution for molecular levels, we average the susceptibility over a Lorentzian distribution of the transition frequencies

\[ P_L(x) = \frac{1}{\pi} \frac{\sigma_x}{(x - \langle x \rangle)^2 + \sigma_x^2}, \]  

(1C)

where \( \langle x \rangle \) and \( \sigma_x \) are the mean value and the standard deviation of the random variable \( x \). The benefit of this approach is that we will get exact analytical averages, which will allow us to gain insight into the system dynamics in simple terms, and help us make meaningful choices of parameters. We show numerically that the results of this procedure compare well with a Gaussian average (see Fig. 6b in the main text).

The proposed technique is based on the observation that there is a class of functions, for which averaging over a Lorentzian distribution can be done instantly. Consider a function of a complex variable \( f(z = \Delta + i\gamma) \), where \( \Delta \) and \( \gamma \) are real quantities; in what follows they will represent, respectively, a random detuning, over which we average, and a constant homogeneous linewidth. We impose two restrictions onto the function \( f(z) \): (1) it must decay faster than \( z \) for \( |z| \to \infty \), and (2) it must not have poles in the upper half-plane. Then averaging of \( f(z) \) over a Lorentzian distribution (C1) \( P_L(\Delta) \) of the detunings with the mean value \( \langle \Delta \rangle \) and a standard deviation \( \sigma \) writes:

\[ \langle f(\Delta + i\gamma) \rangle_{\Delta} = \int_{-\infty}^{\infty} d\Delta \ f(\Delta + i\gamma) \frac{1}{\pi} \frac{\sigma}{(\Delta - \langle \Delta \rangle)^2 + \sigma^2} = \]
\[ = \int_{-\infty}^{\infty} d\Delta \ f(\Delta + i\gamma) \frac{1}{2\pi i} \left[ \frac{1}{\Delta - \langle \Delta \rangle + i\sigma} - \frac{1}{\Delta - \langle \Delta \rangle - i\sigma} \right]. \]  

(1C)
We can calculate this integral by closing the integration contour through the upper half-plane of the complex plane. By assumption (1) the integral over the half-circle vanishes, and by assumption (2) the function $f(z)$ has no poles inside the chosen integration contour, so that the only pole that contributes to the integral is $\Delta = \langle \Delta \rangle + i\sigma$ originating from the Lorentzian distribution. The second integral, $\int_{-\infty}^{\infty} d\Delta \ f(\Delta + i\gamma)[\Delta - (\langle \Delta \rangle - i\sigma)]^{-1}/2\pi i$, thus vanishes. By calculating the residue at the only pole $\Delta = \langle \Delta \rangle + i\sigma$ in the first integral and applying the Cauchy theorem, we get:

$$\langle f(\Delta + i\gamma) \rangle_{\Delta} = f(\langle \Delta \rangle + i[\gamma + \sigma]).$$

(C3)

In summary, as long as the function $f(\Delta + i\gamma)$ satisfies the two criteria listed above, averaging over a Lorentzian distribution consists in replacing the real part of the argument of the function $f(\Delta + i\gamma)$ by its mean value, $\langle \Delta \rangle$, and its imaginary part $\gamma$ by $\gamma + \sigma$, where $\sigma$ is the width of the Lorentzian distribution.

Next, we notice that any susceptibility, being considered as a function of complex frequency, must satisfy the criteria (1) and (2), since they are the same requirements that are imposed on all the material functions which satisfy the Kramers-Kronig relations. Hence, averaging the susceptibility over Lorentzian distribution of the transition frequencies (or the detunings) with the inhomogeneous broadening $\sigma$ can be done by the replacement (C3). The result will be the susceptibility with the fluctuating detuning replaced by its average value, and the homogeneous linewidth $\gamma$ replaced by a sum of the homogeneous and inhomogeneous broadenings, denoted as $\Sigma = \gamma + \sigma$.

Finally, in our multi-level situation, we have to average the susceptibility over more than one Lorentzian distribution. To do this, we repeat this procedure subsequently for each of the disordered transitions. After each averaging we get a new material function that must satisfy the Kramers-Kronig relations, and, consequently, the criteria (1) and (2), so that we can keep repeating this procedure until all the averagings are completed. The result can be symbolically written as (here $\Sigma_i = \gamma_i + \sigma_i$):

$$\langle \chi(\Delta_1 + i\gamma_1, \ldots, \Delta_N + i\gamma_N) \rangle_{\Delta_1, \ldots, \Delta_N} = \chi(\langle \Delta_1 \rangle + i\Sigma_1, \ldots, \langle \Delta_N \rangle + i\Sigma_N).$$

(C4)