Electrochemically Switchable Multimode Strong Coupling in Plasmonic Nanocavities

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ABSTRACT: The strong-coupling interaction between quantum emitters and cavities provides the archetypical platform for fundamental quantum electrodynamics. Here we show that methylene blue (MB) molecules interact coherently with subwavelength plasmonic nanocavity modes at room temperature. Experimental results show that the strong coupling can be switched on and off reversibly when MB molecules undergo redox reactions which transform them to leuco-methylene blue molecules. In simulations we demonstrate the strong coupling between the second excited plasmonic cavity mode and resonant emitters. However, we also show that other detuned modes simultaneously couple efficiently to the molecular transitions, creating unusual cascades of mode spectral shifts and polariton formation. This is possible due to the relatively large plasmonic particle size resulting in reduced mode splittings. The results open significant potential for device applications utilizing active control of strong coupling.

KEYWORDS: multimode strong coupling, strong coupling control, plasmonic nanocavities, polariton formation

The strong coupling regime is central to cavity quantum electrodynamics (cQED), in which the intricate link between quantum emitters and their optical environment leads to the formation of hybrid states that no longer retain their separate identities of light and matter. In this strong coupling regime, the hybrid light-matter states (polaritons) are separated in energy by the vacuum Rabi splitting \( \Omega_R \). Since its discovery for many ultracold atoms in 1989\(^1\) and later for a single atom using a high-finesse optical cavity in 1992,\(^2\) most investigations of strong coupling have aimed at modifying the optical properties, enabling applications such as photon-based quantum information and low-threshold lasing in semiconductors.\(^3\) Through recent advances in nanofabrication,\(^7,8\) plasmonic nanocavities with ultralow cavity mode volumes can confine light into nanometre-sized regions far below the diffraction limit. This provides large coupling strengths, \( \Omega_R \propto 1/\sqrt{V} \), making them a promising approach to access strong coupling at room temperature, at the few- or even single-molecule level.\(^9\) While the rapid evolution of nanoplasmic technologies has enabled a plethora of applications, ranging from single-molecule spectroscopy to solar cells and photothermal cancer treatment,\(^10\)–\(^12\) little information on the available polariton modes and how strong coupling can be dynamically controlled in nanoplasmic environments has been available.

Recently, the selective manipulation of matter by strong coupling and control of interaction strengths from weak to strong coupling has been investigated\(^13\)–\(^15\) with the use of tip-enhanced excitation. Active control of the coupling strength has been demonstrated with photoswitchable molecules,\(^16\)–\(^19\) polarization-dependent coupling between J-aggregates and gold dimers,\(^20\)\(^,\)\(^21\)\(^,\) and electrostatic gating and thermal tuning of monolayer transition metal dichalcogenides.\(^22\)–\(^26\) However, it is challenging to align emitters precisely within a cavity, limiting demonstrations to low yields.\(^27\)

Alternative pathways have emerged that leverage chemistry and material science to guide cQED into new directions and achieve active control of strong coupling easily. In ref 28, Pietron et al. have demonstrated electrochemically controlled switching of strong coupling between molecular vibrational modes and Fabry–Perot cavity modes in the far-infrared spectral range; in particular, the coupling strength was altered through a redox reaction between benzoquinone and dihydroquinone. Naturally, achieving such dynamic switching for electronic transitions interacting with optical-frequency cavity modes would be highly desirable for emerging quantum optoelectronic applications. Our present study reveals that electrochemically triggered redox reactions of methylene blue (MB) molecules can be used to switch on and off strong...
plasmon-exciton interaction in ultracompact, individual nano-
cavities featuring 1 nm nanogaps. Our work accesses a regime
in which a very limited number of molecules (<100) are highly
coupled to the cavity and allow very short switching times
compared to bulk approaches with million-fold larger-volume
Fabry−Perot cavities. Reversible switching is demonstrated
without altering the geometry of the nanocavities. The
sequence of simultaneous multimode plasmonic coupling in
the system is explored using time-domain simulations,
demonstrating the formation of several pairs of polariton states.

The strong coupling is realized using an established model
system consisting of a nanoparticle-on-mirror (NPoM)
nanocavity hosting a layer of optically active methylene blue
(MB) molecules (Figure 1a). The ratio of MB: cucurbit[7]uril (CB)
(1:1) is set by mixing 1 mM solutions of both molecules in
sample preparation, to ensure each MB is coupled with one
CB (see Supporting Information (SI)). The switching of strong
coupling is achieved using an in situ spectro-electrochemical
cell.

In the experiment, automated tracking microscopy is used to
detect dark-field scattering spectra of hundreds of individual
NPoM constructs across the sample surface, where regions of
sparsely distributed NPoMs can be selected to ensure that only
one construct is within the collection area (see Figure S3). In
addition, we perform real-time spectro-electrochemical meas-

Figure 1. Experimental demonstration. (a) Schematic of the NPoM geometry with cucurbit[7]uril (CB) + methylene blue (MB) molecules in
the nanogap layer (purple). The Au nanoparticle has radius \( r = 40 \text{ nm} \) and a facet width of \( w = 20 \text{ nm} \), while the nanogap region has a height of \( d = 1 \text{ nm} \) and a refractive index of 1.4 (CB). The NPoM is illuminated obliquely at a large angle of incidence. Each CB (purple) molecule hosts a single
MB (blue)/leuco-methylene blue (LMB, colorless) molecule ensuring vertical orientation. The circuit is used to perform the reversible redox
reactions from MB (transition at 650 nm) to LMB (transition at 256 nm) by adding/removing two electrons. Chronoamperometry scans are
between +0.5 and −0.5 V, as determined from cyclic voltammetry (Figure S2). Working, counter and reference electrodes are Au the substrate, Pt
mesh, and Ag/AgCl (3 M KCl), respectively. The supporting electrolyte is 0.1 M phosphate buffer solution (PBS) to maintain constant pH 7
during redox. (b) Applied potential (blue) and current density (black) over 30 s, as switch. (c) Evolution of the scattering spectra during the redox
reaction time interval in (b). (d) Evolution of the photoluminescence (PL) spectra during the same time interval.

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measurements of the photoluminescence (PL) yield by applying an electrical bias to the gold substrate while irradiating an individual NPoM with continuous-wave laser light at 532 nm. The current density and potential over a time interval of 30 s are plotted in Figure 1c, where the potential is switched between +0.5 V and −0.5 V (vs Ag/AgCl) every 10 s. This triggers the redox conversion of MB to LMB and back to MB as indicated. The reaction is electrochemically controlled by transferring electrons from the working electrode to the reactant in the nanogap over <1 nm distances (Figure 1b). The simultaneous scattering and PL spectra over the same time interval (Figure 1d, e) clearly identify that both signals change when the molecules are switched. From $t = 10$−20 s, the scattering spectrum of LMB is brighter than MB, while its PL spectrum is dark, because it shifts to an entirely different spectral region and is not excited.

The unusual situation here is that strong coupling is set up not with the fundamental mode but with the higher-order mode $l_z$. This provides an intriguing opportunity to study the additional interplay with further-detuned modes. Since the splitting between plasmonic cavity modes is smaller for larger nanoparticles, $l_z$ does not dominate the spectrum. To unravel this interplay, we first characterize the NPoM cavity and MB coupling to $l_1$, in numerical simulations, adding extra damping to the permittivity of the gold, as detailed in the Methods section. The NPoM system in Figure 1a is first simulated without any active medium to determine the optical field profile as a function of the in-plane position $x$ for the 500−900 nm wavelength range of interest (Figure 2a). We find the two dominant plasmonic modes $l_1$ and $l_z$ at 840 and 615 nm, respectively (dashed lines). To confirm their mode character the corresponding mode profiles are extracted (Figure 2b).

The expected single peak for the lowest energy mode $l_1$ (green) is seen at 840 nm, while the energetically higher mode $l_z$ at 615 nm (red) shows two nodes, which stem from a single nodal circle in the $xy$ plane (see Figure 3 below). The slight asymmetry observed in the mode profiles of Figures 2b and 3c stems from the choice of incident light direction used in the simulation. The light is incident from the left side of the NPoM geometry with polarization along the $z$-axis into the positive $x$-direction, which breaks the symmetry between $x$- and $y$-directions. While the probing light is incident obliquely in the experiments, our use of laterally incident light in the numerical simulations ensures efficient excitation of the NPoM cavity modes, which bear a largely vertical intracavity field polarization. This choice also ensures the excitation of the $E_z$-polarized lowest $l_1$ mode.

Since the absorption line of the MB molecules is affected by their dielectric environment, its exact resonance position is not possible to extract from nanogap experiments. Therefore, we model the MB molecules as two-level quantum emitters in our simulations and vary the detuning between the emitters and the cavity mode $l_z$ by tuning the transition wavelength of the emitters $\lambda_e$ from 570 to 680 nm across the $l_z$ resonance at 615 nm.

We choose the density of emitters such that there are on average 50 emitters under the NPoM facet, each with a dipole moment of $\mu = 4 \text{ D}$ and dephasing rate of $5 \times 10^{13} \text{ rad/s}$ (HWHM $\Gamma$ of 33 meV). The corresponding results for the scattering spectra are shown in Figure 2c. The clear anticrossing behavior forming two polariton branches around vanishing detuning at $\lambda_e = 615 \text{ nm}$ indicates the occurrence of strong coupling.

From basic models of multimitter strong coupling, it is known that the coupling strength scales as $g \propto \mu\sqrt{N}$ for $N$ emitters in resonance with a single plasmonic mode. We conduct a systematic study to investigate the impact of the number of emitters on the coupling strength in the resonant case, since it is difficult to determine the average number of MB molecules in each NPoM cavity in experiment. Scattering spectra are simulated for a range of emitter densities from 0.01 to 0.3 molecules/nm$^3$, corresponding to average molecule numbers under the facet from 10 to 120 which is a reasonable range for the experimental molecular density under typical NPoM facets. The polariton peak positions (Figure 2d, red circles) extracted for each emitter density show the peak splitting exhibits the expected square root dependence on the number of emitters (red solid line).

Larger NPoM systems ($r > 30 \text{ nm}$) have sufficiently dense plasmonic modes that many NPoM modes have relatively small detunings to the MB transition. Consequently, the MB molecules can couple to multiple plasmonic modes simultaneously and multimode strong coupling emerges for sufficiently large coupling strengths. Surprisingly, we find that the polaritons arise independently from each plasmonic mode, manifesting in the formation of multiple pairs of polaritons observed as peaks in the scattering spectrum.

To systematically study multimode coupling, we simulate the system when varying the dipole moment of emitters from 0 to 11 D while fixing the MB transition at 650 nm with 50 molecules under the NPoM facet. To make evident the multimode coupling behavior in simulations, the extra damping of gold is back to normal and the dephasing rate of...
The calculated scattering spectra (Figure 3a) show that coupling between each plasmonic mode and the emitters always exists and is proportional to the dipole moment independent of detuning. Although the effective coupling is significantly reduced for strongly detuned plasmonic modes, the splitting effect from polariton formation leads to energy shifts for all modes. As these shifts overcome the line broadening for sufficiently large coupling, they can be distinguished individually. The scattering spectra for three different dipole moments (Figure 3b, μ = 0, 4, 10 D), marked with dashed lines in Figure 3a, identify several peaks for larger dipole moments. By μ = 10 D, 7 peaks (including the perturbed l1) are identified, while the shift of polariton peaks between 600−700 nm (Figure 3a) is near-linear with dipole moment, as expected.

Figure 3. Simulation of multimode strong coupling. (a) Scattering spectra with varying magnitude of dipole moment from μ = 0 to 11 D. (b) Scattering spectra for dipole moments of μ = 0, 4, and 10 D from bottom to top as marked by dashed lines in (a). (c) Ex mode profiles of each peak in the scattering spectrum at μ = 10 D at indicated wavelengths. Top row: field in xy plane across the middle of the nanogap (red corresponding to positive, blue to negative Ez). Bottom row: 1 D cuts of Ez along the x-axis at y = 0 nm. Pairs of polaritons can be identified by the number of nodes and are indicated by index 1−4; +/− signs indicate the upper and lower polaritons, respectively.

Figure 4. Comparison between scattering spectra in simulation and experiment. (a−c) Simulation and (d−f) experiment, for modes around (a, b, d, e) l2 and (c, f) l1, where gap molecules are (a,d) LMB (uncoupled, orange) or (b, e) MB (coupled, blue). The dashed black lines indicate Gaussian fits, while solid black lines indicate the component peaks. The spectral position of l2 (dashed vertical lines) is indicated in (b, e). The polariton peaks 2± have a combined width of 133 meV in simulation and (181 ± 5) meV in experiment and a splitting of 145 meV in simulation and (172 ± 5) meV in experiment.
More careful analysis identifies pairs of polaritons by comparing the spatial distributions of optical fields at all peaks in the scattering spectrum. At $\mu = 10 \text{ D}$, spatial distributions of $E_z(x, y)$ through the middle of the nanogap (upper row, Figure 3c) and of $E_z(x)$ at $y = 0 \text{ nm}$ (bottom row) are extracted for the seven labeled peaks in the scattering spectrum. Each NPoM mode couples to the emitters with a corresponding coupling strength and forms a lower polariton ($-$) and an upper polariton ($+$), with these polariton pairs indexed 1–3 (see labels). At $\mu = 0 \text{ D}$ these pairs are degenerate at the respective bare plasmonic modes $l_{1,3}$. Extracting the mode positions shows their unusual hierarchy of splitting around the electronic resonance (see SI Figures S5 and S7).

Taking the strong coupling between mode $l_1$ and the emitters as an example, the hybrid states $1^+ \text{ and } 1^-$ are formed at spectral positions 635 and 957 nm. We can pair these peaks through their spatial distributions of $E_y$ as both have similar structures with a single maximum (red circular shape) and no nodes (Figure 3c). This characteristic spatial distribution is inherited from mode $l_1$ which shows a single antinode in the mode profile. The same procedure can be carried out for the next higher modes leading to the full polariton labeling (Figure 3b, c). The mode profiles become clearer and more pronounced with larger coupling strength. We emphasize how the polariton mode spatial profiles in the multimode strong coupling situation (Figure 3) become nontrivial superpositions of light and matter. We see that the polariton splittings increase with different rates depending on the effective coupling strengths. This means that, for very large dipole moments, different polariton resonances can spectrally intersect, form new hybrid polaritons, and develop additional anticrossings. This is demonstrated in SI Figure S7 depicting a larger dipole moment scan for the parameters from Figure 3a.

We now return to the measured scattering spectra with MB and LMB molecules and compare them with our theoretical predications for coupled and uncoupled nanocavity systems. Our goal is to identify spectral features demonstrating the switching of strong coupling in the system. Comparing the scattering spectra of uncoupled nanocavities in simulation and experiment (Figure 4a, d) shows that the spectral position of the mode $l_2$ is well matched at $\sim 615 \text{ nm}$, although the broadening is significantly stronger in experiment. We model the same NPoM system with 50 emitters under the facet, a dipole moment of 4 D, transition wavelength of 650 nm, and total dephasing rate of $1 \times 10^{14} \text{ rad/s} \ (\Gamma = 66 \text{ meV})$. For the emitter-coupled nanocavities (Figure 4b, e), a shoulder appears on the high-wavelength side of the $l_2$ peak, arising from the MB resonance. To quantify all relevant spectral positions, we fit the spectra carefully with Gaussian lineshapes (dashed lines), with the two closest individual peaks plotted (solid curves, Figures 4b, e). Two dependencies are noted: (i) an additional peak appears around 650 nm from the MB resonance, and (ii) both peaks are shifted away from their uncoupled position (dashed vertical lines). Feature (ii) demonstrates the creation of polariton states and the appearance of a near-strong-coupling regime. The splitting of the polariton peaks in the experimental data (Figure 4e) is harder to identify from the scattering spectra collected in an electrochemical cell as our experiments in glass cells with water restrict high-angle light collection. We thus also perform experiments outside the electrochemical cell for 50 different NPoMs (Figure S1). These indeed show a clear difference with and without coupling between the MB molecules and the $l_1$ plasmon mode. It is important to note that we not only observe a change at the $l_2$ mode but also a spectral shift of the $l_1$ mode, as correctly predicted by our simulations. Further we clarify that while Figure 4e does not quite resolve clear strong coupling, in correspondence with our simulations modeling the scattering spectra, we find two polariton peaks at the wavelengths predicted by simulation. We also note in Figure S1 that the different nanoparticles are neither identical in diameter nor identically shaped, which makes perfect agreement between simulations and experiment unviable. However, our simulations predict the correct physical behavior assuming reasonable system parameters.

Examining now the modes around $l_1$ (Figure 4c, f) there is only a small shift from the uncoupled (orange) to the coupled (blue) mode in the simulation. This is because the dipole moment of MB molecules (4 D) provides insufficient coupling strength for realizing strong coupling with $l_1$ in the presence of significant detuning. The larger shift of $l_1$ in experiment (Figure 4f) suggests additional effects from the redox process, such as changing the refractive index around the nanoparticle.\(^{38,39}\) We note the experimental scattering is also broader and weaker than theory predicts, suggesting effects from disorder and/or inhomogeneity.

Strong coupling has been observed in plasmonic nanocavities at room temperature and several experiments have proposed to investigate tuning of the coupling strength. Photoisomerization can also switch electronic resonances, but is typically slow (on time scales of tens to hundreds of minutes). By contrast, our observations show electrochemical control of strong coupling on subsecond time scales. Because the redox process requires electron transfer over subnanometre length scales, our scheme can potentially reach or even exceed GHz operation rates. Redox reactions of MB molecules allow the reversible and nondestructive, on–off switching of the strong coupling without altering the geometry of the nanocavities.

We find that multimode strong coupling in plasmonic systems retains the nanocavity mode symmetry, providing overlapping pairs of polariton modes as the dipole strength increases. This unusual situation comes from the low mode volumes in such plasmonic nanocavities, but also the specific spectrum of their resonance frequencies. Simulations verify that the coupling strength scales as $\mu \sqrt{N}$. We anticipate that electrically switchable strong coupling will facilitate many developments in the fields of quantum chemistry, nonlinear optics, and molecular quantum optics.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.3c03814. (1) Experimental methods: sample design, sample preparation, Spectro-electrochemical cells, Optical measurements, Cyclic voltammetry (CV), NPoM density, MB absorption spectrum; (2) Theoretical methods: FDTD method, Two-level Maxwell-Bloch model, Metal–insulator–metal waveguide model, classical coupled-oscillator model, higher-order hybrid states. (PDF)
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Notes
The authors declare no competing financial interest.

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